COMPOSITIONS AND METHODS FOR THE TREATMENT OF IMMUNE RELATED DISEASES

Field of the Invention

The present invention relates to compositions and methods useful for the diagnosis and treatment of immune related diseases.

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Background of the Invention

Immune related and inflammatory diseases are the manifestation or consequence of fairly complex, often multiple interconnected biological pathways which in normal physiology are critical to respond to insult or injury, initiate repair from insult or injury, and mount innate and acquired defense against foreign organisms. Disease or pathology occurs when these normal physiological pathways cause additional insult or injury either as directly related to the intensity of the response, as a consequence of abnormal regulation or excessive stimulation, as a reaction to self, or as a combination of these.

Though the genesis of these diseases often involves multistep pathways and often multiple different biological systems/pathways, intervention at critical points in one or more of these pathways can have an ameliorative or therapeutic effect. Therapeutic intervention can occur by either antagonism of a detrimental process/pathway or stimulation of a beneficial process/pathway.

Many immune related diseases are known and have been extensively studied. Such diseases include immune-mediated inflammatory diseases, non-immune-mediated inflammatory diseases, infectious diseases, immunodeficiency diseases, neoplasia, etc.

T lymphocytes (T cells) are an important component of a mammalian immune response. T cells recognize antigens which are associated with a self-molecule encoded by genes within the major histocompatibility complex (MHC). The antigen may be displayed together with MHC molecules on the surface of antigen presenting cells, virus infected cells, cancer cells, grafts, etc. The T cell system eliminates these altered cells which pose a health threat to the host mammal. T cells include helper T cells and cytotoxic T cells. Helper T cells proliferate extensively following recognition of an antigen -MHC complex on an antigen presenting cell. Helper T cells also secrete a variety of cytokines, i.e., lymphokines, which play a central role in the activation of B cells, cytotoxic T cells and a variety of other cells which participate in the immune response.

Immune related diseases could be treated by suppressing the immune response. Using neutralizing antibodies that inhibit molecules having immune stimulatory activity would be beneficial in the treatment of immune-mediated and inflammatory diseases. Molecules which inhibit the immune response can be utilized (proteins directly or via the use of antibody agonists) to inhibit the immune response and thus ameliorate immune related disease.

CD4+ T cells are known to be important regulators of inflammation. Herein, CD4+ T cells were activated and the profile of genes differentially expressed upon activation was analyzed. As such, the activation specific genes may be potential therapeutic targets. *In vivo* co-stimulation is necessary for a productive immune proliferative response. The list of costimulatory molecules is quite extensive and it is still unclear just which co-stimulatory molecules play critical roles in different types and stages of

inflammation. In this application the focus is on genes which are specifically upregulated or downregulated by stimulation with anti-CD3/ICAM, or anti-CD3/anti-CD28 and may be useful in targeting inflammatory processes which are associated with these different molecules.

Despite the above identified advances in T cell research, there is a great need for additional diagnostic and therapeutic agents capable of detecting the presence of a T cell mediated disorders in a mammal and for effectively reducing these disorders. Accordingly, it is an objective of the present invention to identify polypeptides that are overexpressed in activated T cells as compared to resting T cells, and to use those polypeptides, and their encoding nucleic acids, to produce compositions of matter useful in the therapeutic treatment and diagnostic detection of T cell mediated disorders in mammals.

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Summary of the Invention

A. Embodiments

The present invention concerns compositions and methods useful for the diagnosis and treatment of immune related disease in mammals, including humans. The present invention is based on the identification of proteins (including agonist and antagonist antibodies) which are a result of stimulation of the immune response in mammals. Immune related diseases can be treated by suppressing or enhancing the immune response. Molecules that enhance the immune response stimulate or potentiate the immune response to an antigen. Molecules which stimulate the immune response can be used therapeutically where enhancement of the immune response would be beneficial. Alternatively, molecules that suppress the immune response attenuate or reduce the immune response to an antigen (e.g., neutralizing antibodies) can be used therapeutically where attenuation of the immune response would be beneficial (e.g., inflammation). Accordingly, the PRO polypeptides, agonists and antagonists thereof are also useful to prepare medicines and medicaments for the treatment of immune-related and inflammatory diseases. In a specific aspect, such medicines and medicaments comprise a therapeutically effective amount of a PRO polypeptide, agonist or antagonist thereof with a pharmaceutically acceptable carrier. Preferably, the admixture is sterile.

In a further embodiment, the invention concerns a method of identifying agonists or antagonists to a PRO polypeptide which comprises contacting the PRO polypeptide with a candidate molecule and monitoring a biological activity mediated by said PRO polypeptide. Preferably, the PRO polypeptide is a native sequence PRO polypeptide. In a specific aspect, the PRO agonist or antagonist is an anti-PRO antibody.

In another embodiment, the invention concerns a composition of matter comprising a PRO polypeptide or an agonist or antagonist antibody which binds the polypeptide in admixture with a carrier or excipient. In one aspect, the composition comprises a therapeutically effective amount of the polypeptide or antibody. In another aspect, when the composition comprises an immune stimulating molecule, the composition is useful for: (a) increasing infiltration of inflammatory cells into a tissue of a mammal in need thereof, (b) stimulating or enhancing an immune response in a mammal in need thereof, (c) increasing the proliferation of T-lymphocytes in a mammal in need thereof in response to an antigen, (d) stimulating the activity of T-lymphocytes or (e) increasing the vascular permeability. In a further aspect, when the composition comprises an immune inhibiting molecule, the composition is useful for: (a) decreasing infiltration of inflammatory cells into a tissue of a mammal in need thereof, (b) inhibiting or reducing an

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immune response in a mammal in need thereof, (c) decreasing the activity of T-lymphocytes or (d) decreasing the proliferation of T-lymphocytes in a mammal in need thereof in response to an antigen. In another aspect, the composition comprises a further active ingredient, which may, for example, be a further antibody or a cytotoxic or chemotherapeutic agent. Preferably, the composition is sterile.

In another embodiment, the invention concerns a method of treating an immune related disorder in a mammal in need thereof, comprising administering to the mammal an effective amount of a PRO polypeptide, an agonist thereof, or an antagonist thereto. In a preferred aspect, the immune related disorder is selected from the group consisting of: systemic lupus erythematosis, rheumatoid arthritis, osteoarthritis, juvenile chronic arthritis, spondyloarthropathies, systemic sclerosis, idiopathic inflammatory myopathies, Sjögren's syndrome, systemic vasculitis, sarcoidosis, autoimmune hemolytic anemia, autoimmune thrombocytopenia, thyroiditis, diabetes mellitus, immune-mediated renal disease, demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy, hepatobiliary diseases such as infectious, autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis, inflammatory bowel disease, gluten-sensitive enteropathy, and Whipple's disease, autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis, allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria, immunologic diseases of the lung such as eosinophilic pneumonias, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis, transplantation associated diseases including graft rejection and graft -versus-host-disease.

In another embodiment, the invention provides an antibody which specifically binds to any of the above or below described polypeptides. Optionally, the antibody is a monoclonal antibody, humanized antibody, antibody fragment or single-chain antibody. In one aspect, the present invention concerns an isolated antibody which binds a PRO polypeptide. In another aspect, the antibody mimics the activity of a PRO polypeptide (an agonist antibody) or conversely the antibody inhibits or neutralizes the activity of a PRO polypeptide (an antagonist antibody). In another aspect, the antibody is a monoclonal antibody, which preferably has nonhuman complementarity determining region (CDR) residues and human framework region (FR) residues. The antibody may be labeled and may be immobilized on a solid support. In a further aspect, the antibody is an antibody fragment, a monoclonal antibody, a single-chain antibody, or an anti-idiotypic antibody.

In yet another embodiment, the present invention provides a composition comprising an anti-PRO antibody in admixture with a pharmaceutically acceptable carrier. In one aspect, the composition comprises a therapeutically effective amount of the antibody. Preferably, the composition is sterile. The composition may be administered in the form of a liquid pharmaceutical formulation, which may be preserved to achieve extended storage stability. Alternatively, the antibody is a monoclonal antibody, an antibody fragment, a humanized antibody, or a single-chain antibody.

In a further embodiment, the invention concerns an article of manufacture, comprising:

- (a) a composition of matter comprising a PRO polypeptide or agonist or antagonist thereof;
- (b) a container containing said composition; and

(c) a label affixed to said container, or a package insert included in said container referring to the use of said PRO polypeptide or agonist or antagonist thereof in the treatment of an immune related disease. The composition may comprise a therapeutically effective amount of the PRO polypeptide or the agonist or antagonist thereof.

In yet another embodiment, the present invention concerns a method of diagnosing an immune related disease in a mammal, comprising detecting the level of expression of a gene encoding a PRO polypeptide (a) in a test sample of tissue cells obtained from the mammal, and (b) in a control sample of known normal tissue cells of the same cell type, wherein a higher or lower expression level in the test sample as compared to the control sample indicates the presence of immune related disease in the mammal from which the test tissue cells were obtained.

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In another embodiment, the present invention concerns a method of diagnosing an immune disease in a mammal, comprising (a) contacting an anti-PRO antibody with a test sample of tissue cells obtained from the mammal, and (b) detecting the formation of a complex between the antibody and a PRO polypeptide, in the test sample; wherein the formation of said complex is indicative of the presence or absence of said disease. The detection may be qualitative or quantitative, and may be performed in comparison with monitoring the complex formation in a control sample of known normal tissue cells of the same cell type. A larger quantity of complexes formed in the test sample indicates the presence or absence of an immune disease in the mammal from which the test tissue cells were obtained. The antibody preferably carries a detectable label. Complex formation can be monitored, for example, by light microscopy, flow cytometry, fluorimetry, or other techniques known in the art. The test sample is usually obtained from an individual suspected of having a deficiency or abnormality of the immune system.

In another embodiment, the invention provides a method for determining the presence of a PRO polypeptide in a sample comprising exposing a test sample of cells suspected of containing the PRO polypeptide to an anti-PRO antibody and determining the binding of said antibody to said cell sample. In a specific aspect, the sample comprises a cell suspected of containing the PRO polypeptide and the antibody binds to the cell. The antibody is preferably detectably labeled and/or bound to a solid support.

In another embodiment, the present invention concerns an immune-related disease diagnostic kit, comprising an anti-PRO antibody and a carrier in suitable packaging. The kit preferably contains instructions for using the antibody to detect the presence of the PRO polypeptide. Preferably the carrier is pharmaceutically acceptable.

In another embodiment, the present invention concerns a diagnostic kit, containing an anti-PRO antibody in suitable packaging. The kit preferably contains instructions for using the antibody to detect the PRO polypeptide.

In another embodiment, the invention provides a method of diagnosing an immune-related disease in a mammal which comprises detecting the presence or absence or a PRO polypeptide in a test sample of tissue cells obtained from said mammal, wherein the presence or absence of the PRO polypeptide in said test sample is indicative of the presence of an immune-related disease in said mammal.

In another embodiment, the present invention concerns a method for identifying an agonist of a PRO polypeptide comprising:

(a) contacting cells and a test compound to be screened under conditions suitable for the induction of a cellular response normally induced by a PRO polypeptide; and

(b) determining the induction of said cellular response to determine if the test compound is an effective agonist, wherein the induction of said cellular response is indicative of said test compound being an effective agonist.

In another embodiment, the invention concerns a method for identifying a compound capable of inhibiting the activity of a PRO polypeptide comprising contacting a candidate compound with a PRO polypeptide under conditions and for a time sufficient to allow these two components to interact and determining whether the activity of the PRO polypeptide is inhibited. In a specific aspect, either the candidate compound or the PRO polypeptide is immobilized on a solid support. In another aspect, the non-immobilized component carries a detectable label. In a preferred aspect, this method comprises the steps of:

- (a) contacting cells and a test compound to be screened in the presence of a PRO polypeptide under conditions suitable for the induction of a cellular response normally induced by a PRO polypeptide; and
 - (b) determining the induction of said cellular response to determine if the test compound is an effective antagonist.

In another embodiment, the invention provides a method for identifying a compound that inhibits the expression of a PRO polypeptide in cells that normally express the polypeptide, wherein the method comprises contacting the cells with a test compound and determining whether the expression of the PRO polypeptide is inhibited. In a preferred aspect, this method comprises the steps of:

- (a) contacting cells and a test compound to be screened under conditions suitable for allowing expression of the PRO polypeptide; and
 - (b) determining the inhibition of expression of said polypeptide.

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In yet another embodiment, the present invention concerns a method for treating an immune-related disorder in a mammal that suffers therefrom comprising administering to the mammal a nucleic acid molecule that codes for either (a) a PRO polypeptide, (b) an agonist of a PRO polypeptide or (c) an antagonist of a PRO polypeptide, wherein said agonist or antagonist may be an anti-PRO antibody. In a preferred embodiment, the mammal is human. In another preferred embodiment, the nucleic acid is administered via ex vivo gene therapy. In a further preferred embodiment, the nucleic acid is comprised within a vector, more preferably an adenoviral, adeno-associated viral, lentiviral or retroviral vector.

In yet another aspect, the invention provides a recombinant viral particle comprising a viral vector consisting essentially of a promoter, nucleic acid encoding (a) a PRO polypeptide, (b) an agonist polypeptide of a PRO polypeptide, or (c) an antagonist polypeptide of a PRO polypeptide, and a signal sequence for cellular secretion of the polypeptide, wherein the viral vector is in association with viral structural proteins. Preferably, the signal sequence is from a mammal, such as from a native PRO polypeptide.

In a still further embodiment, the invention concerns an ex vivo producer cell comprising a nucleic acid construct that expresses retroviral structural proteins and also comprises a retroviral vector consisting essentially of a promoter, nucleic acid encoding (a) a PRO polypeptide, (b) an agonist polypeptide of a PRO polypeptide or (c) an antagonist polypeptide of a PRO polypeptide, and a signal sequence for cellular secretion of the polypeptide, wherein said producer cell packages the retroviral vector in association with the structural proteins to produce recombinant retroviral particles.

In a still further embodiment, the invention provides a method of increasing the activity of T-lymphocytes in a mammal comprising administering to said mammal (a) a PRO polypeptide, (b) an agonist of a PRO polypeptide, or (c) an antagonist of a PRO polypeptide, wherein the activity of T-lymphocytes in the mammal is increased.

In a still further embodiment, the invention provides a method of decreasing the activity of T-lymphocytes in a mammal comprising administering to said mammal (a) a PRO polypeptide, (b) an agonist of a PRO polypeptide, or (c) an antagonist of a PRO polypeptide, wherein the activity of T-lymphocytes in the mammal is decreased.

In a still further embodiment, the invention provides a method of increasing the proliferation of T-lymphocytes in a mammal comprising administering to said mammal (a) a PRO polypeptide, (b) an agonist of a PRO polypeptide, or (c) an antagonist of a PRO polypeptide, wherein the proliferation of T-lymphocytes in the mammal is increased.

In a still further embodiment, the invention provides a method of decreasing the proliferation of T-lymphocytes in a mammal comprising administering to said mammal (a) a PRO polypeptide, (b) an agonist of a PRO polypeptide, or (c) an antagonist of a PRO polypeptide, wherein the proliferation of T-lymphocytes in the mammal is decreased.

B. Additional Embodiments

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In other embodiments of the present invention, the invention provides vectors comprising DNA encoding any of the herein described polypeptides. Host cell comprising any such vector are also provided. By way of example, the host cells may be CHO cells, *E. coli*, or yeast. A process for producing any of the herein described polypeptides is further provided and comprises culturing host cells under conditions suitable for expression of the desired polypeptide and recovering the desired polypeptide from the cell culture.

In other embodiments, the invention provides chimeric molecules comprising any of the herein described polypeptides fused to a heterologous polypeptide or amino acid sequence. Example of such chimeric molecules comprise any of the herein described polypeptides fused to an epitope tag sequence or a Fc region of an immunoglobulin.

In another embodiment, the invention provides an antibody which specifically binds to any of the above or below described polypeptides. Optionally, the antibody is a monoclonal antibody, humanized antibody, antibody fragment or single-chain antibody.

In yet other embodiments, the invention provides oligonucleotide probes useful for isolating genomic and cDNA nucleotide sequences or as antisense probes, wherein those probes may be derived from any of the above or below described nucleotide sequences.

In other embodiments, the invention provides an isolated nucleic acid molecule comprising a nucleotide sequence that encodes a PRO polypeptide.

In one aspect, the isolated nucleic acid molecule comprises a nucleotide sequence having at least about 80% nucleic acid sequence identity, alternatively at least about 81% nucleic acid sequence identity, alternatively at least about 83% nucleic acid sequence identity, alternatively at least about 83% nucleic acid sequence identity, alternatively at least about 85% nucleic acid sequence identity, alternatively at least about 85% nucleic acid sequence identity, alternatively at least about 86% nucleic acid sequence identity,

alternatively at least about 87% nucleic acid sequence identity, alternatively at least about 88% nucleic acid sequence identity, alternatively at least about 99% nucleic acid sequence identity, alternatively at least about 90% nucleic acid sequence identity, alternatively at least about 91% nucleic acid sequence identity, alternatively at least about 92% nucleic acid sequence identity, alternatively at least about 93% nucleic acid sequence identity, alternatively at least about 95% nucleic acid sequence identity, alternatively at least about 95% nucleic acid sequence identity, alternatively at least about 96% nucleic acid sequence identity, alternatively at least about 98% nucleic acid sequence identity at least about 98% nucleic acid sequence identity and alternatively at least about 99% nucleic acid sequence identity to (a) a DNA molecule encoding a PRO polypeptide having a full-length amino acid sequence as disclosed herein, an amino acid sequence lacking the signal peptide as disclosed herein, an extracellular domain of a transmembrane protein, with or without the signal peptide, as disclosed herein or any other specifically defined fragment of the full-length amino acid sequence as disclosed herein, or (b) the complement of the DNA molecule of (a).

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In other aspects, the isolated nucleic acid molecule comprises a nucleotide sequence having at least about 80% nucleic acid sequence identity, alternatively at least about 81% nucleic acid sequence identity, alternatively at least about 82% nucleic acid sequence identity, alternatively at least about 83% nucleic acid sequence identity, alternatively at least about 84% nucleic acid sequence identity, alternatively at least about 85% nucleic acid sequence identity, alternatively at least about 86% nucleic acid sequence identity, alternatively at least about 87% nucleic acid sequence identity, alternatively at least about 88% nucleic acid sequence identity, alternatively at least about 89% nucleic acid sequence identity, alternatively at least about 90% nucleic acid sequence identity, alternatively at least about 91% nucleic acid sequence identity, alternatively at least about 92% nucleic acid sequence identity, alternatively at least about 93% nucleic acid sequence identity, alternatively at least about 94% nucleic acid sequence identity, alternatively at least about 95% nucleic acid sequence identity, alternatively at least about 96% nucleic acid sequence identity, alternatively at least about 97% nucleic acid sequence identity, alternatively at least about 98% nucleic acid sequence identity and alternatively at least about 99% nucleic acid sequence identity to (a) a DNA molecule comprising the coding sequence of a full-length PRO polypeptide cDNA as disclosed herein, the coding sequence of a PRO polypeptide lacking the signal peptide as disclosed herein, the coding sequence of an extracellular domain of a transmembrane PRO polypeptide, with or without the signal peptide, as disclosed herein or the coding sequence of any other specifically defined fragment of the full-length amino acid sequence as disclosed herein, or (b) the complement of the DNA molecule of (a).

In a further aspect, the invention concerns an isolated nucleic acid molecule comprising a nucleotide sequence having at least about 80% nucleic acid sequence identity, alternatively at least about 81% nucleic acid sequence identity, alternatively at least about 82% nucleic acid sequence identity, alternatively at least about 83% nucleic acid sequence identity, alternatively at least about 84% nucleic acid sequence identity, alternatively at least about 85% nucleic acid sequence identity, alternatively at least about 86% nucleic acid sequence identity, alternatively at least about 87% nucleic acid sequence identity, alternatively at least about 89% nucleic acid sequence identity, alternatively at least about 89% nucleic acid sequence identity, alternatively at least about 91% nucleic acid sequence identity, alternatively at least about 92% nucleic acid sequence identity, alternatively at least about 94% nucleic acid sequence identity, alternatively at least about 94% nucleic acid sequence identity, alternatively at least about 94% nucleic acid

sequence identity, alternatively at least about 95% nucleic acid sequence identity, alternatively at least about 96% nucleic acid sequence identity, alternatively at least about 97% nucleic acid sequence identity, alternatively at least about 98% nucleic acid sequence identity and alternatively at least about 99% nucleic acid sequence identity to (a) a DNA molecule that encodes the same mature polypeptide encoded by any of the human protein cDNAs as disclosed herein, or (b) the complement of the DNA molecule of (a).

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Another aspect the invention provides an isolated nucleic acid molecule comprising a nucleotide sequence encoding a PRO polypeptide which is either transmembrane domain-deleted or transmembrane domain-inactivated, or is complementary to such encoding nucleotide sequence, wherein the transmembrane domain(s) of such polypeptide are disclosed herein. Therefore, soluble extracellular domains of the herein described PRO polypeptides are contemplated.

Another embodiment is directed to fragments of a PRO polypeptide coding sequence, or the complement thereof, that may find use as, for example, hybridization probes, for encoding fragments of a PRO polypeptide that may optionally encode a polypeptide comprising a binding site for an anti-PRO antibody or as antisense oligonucleotide probes. Such nucleic acid fragments are usually at least about 20 nucleotides in length, alternatively at least about 30 nucleotides in length, alternatively at least about 40 nucleotides in length, alternatively at least about 50 nucleotides in length, alternatively at least about 60 nucleotides in length, alternatively at least about 70 nucleotides in length, alternatively at least about 80 nucleotides in length, alternatively at least about 90 nucleotides in length, alternatively at least about 100 nucleotides in length, alternatively at least about 110 nucleotides in length, alternatively at least about 120 nucleotides in length, alternatively at least about 130 nucleotides in length, alternatively at least about 140 nucleotides in length, alternatively at least about 150 nucleotides in length, alternatively at least about 160 nucleotides in length, alternatively at least about 170 nucleotides in length, alternatively at least about 180 nucleotides in length, alternatively at least about 190 nucleotides in length, alternatively at least about 200 nucleotides in length, alternatively at least about 250 nucleotides in length, alternatively at least about 300 nucleotides in length, alternatively at least about 350 nucleotides in length, alternatively at least about 400 nucleotides in length, alternatively at least about 450 nucleotides in length, alternatively at least about 500 nucleotides in length, alternatively at least about 600 nucleotides in length, alternatively at least about 700 nucleotides in length, alternatively at least about 800 nucleotides in length, alternatively at least about 900 nucleotides in length and alternatively at least about 1000 nucleotides in length, wherein in this context the term "about" means the referenced nucleotide sequence length plus or minus 10% of that referenced length. It is noted that novel fragments of a PRO polypeptide-encoding nucleotide sequence may be determined in a routine manner by aligning the PRO polypeptide-encoding nucleotide sequence with other known nucleotide sequences using any of a number of well known sequence alignment programs and determining which PRO polypeptide-encoding nucleotide sequence fragment(s) are novel. All of such PRO polypeptide-encoding nucleotide sequences are contemplated herein. Also contemplated are the PRO polypeptide fragments encoded by these nucleotide molecule fragments, preferably those PRO polypeptide fragments that comprise a binding site for an anti-PRO antibody.

In another embodiment, the invention provides isolated PRO polypeptide encoded by any of the isolated nucleic acid sequences herein above identified.

In a certain aspect, the invention concerns an isolated PRO polypeptide, comprising an amino acid sequence having at least about 80% amino acid sequence identity, alternatively at least about 81% amino acid sequence identity, alternatively at least about 82% amino acid sequence identity, alternatively at least about 83% amino acid sequence identity, alternatively at least about 84% amino acid sequence identity, alternatively at least about 85% amino acid sequence identity, alternatively at least about 86% amino acid sequence identity, alternatively at least about 87% amino acid sequence identity, alternatively at least about 88% amino acid sequence identity, alternatively at least about 89% amino acid sequence identity, alternatively at least about 90% amino acid sequence identity, alternatively at least about 91% amino acid sequence identity, alternatively at least about 92% amino acid sequence identity, alternatively at least about 93% amino acid sequence identity, alternatively at least about 94% amino acid sequence identity, alternatively at least about 95% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 97% amino acid sequence identity, alternatively at least about 98% amino acid sequence identity and alternatively at least about 99% amino acid sequence identity to a PRO polypeptide having a full-length amino acid sequence as disclosed herein, an amino acid sequence lacking the signal peptide as disclosed herein, an extracellular domain of a transmembrane protein, with or without the signal peptide, as disclosed herein or any other specifically defined fragment of the full-length amino acid sequence as disclosed herein.

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In a further aspect, the invention concerns an isolated PRO polypeptide comprising an amino acid sequence having at least about 80% amino acid sequence identity, alternatively at least about 81% amino acid sequence identity, alternatively at least about 83% amino acid sequence identity, alternatively at least about 83% amino acid sequence identity, alternatively at least about 85% amino acid sequence identity, alternatively at least about 86% amino acid sequence identity, alternatively at least about 86% amino acid sequence identity, alternatively at least about 88% amino acid sequence identity, alternatively at least about 89% amino acid sequence identity, alternatively at least about 91% amino acid sequence identity, alternatively at least about 91% amino acid sequence identity, alternatively at least about 93% amino acid sequence identity, alternatively at least about 93% amino acid sequence identity, alternatively at least about 95% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity and alternatively at least about 99% amino acid sequence identity to an amino acid sequence encoded by any of the human protein cDNAs as disclosed herein.

In a specific aspect, the invention provides an isolated PRO polypeptide without the N-terminal signal sequence and/or the initiating methionine and is encoded by a nucleotide sequence that encodes such an amino acid sequence as herein before described. Processes for producing the same are also herein described, wherein those processes comprise culturing a host cell comprising a vector which comprises the appropriate encoding nucleic acid molecule under conditions suitable for expression of the PRO polypeptide and recovering the PRO polypeptide from the cell culture.

Another aspect the invention provides an isolated PRO polypeptide which is either transmembrane domain-deleted or transmembrane domain-inactivated. Processes for producing the same are also herein described, wherein those processes comprise culturing a host cell comprising a vector which comprises the

appropriate encoding nucleic acid molecule under conditions suitable for expression of the PRO polypeptide and recovering the PRO polypeptide from the cell culture.

In yet another embodiment, the invention concerns agonists and antagonists of a native PRO polypeptide as defined herein. In a particular embodiment, the agonist or antagonist is an anti-PRO antibody or a small molecule.

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In a further embodiment, the invention concerns a method of identifying agonists or antagonists to a PRO polypeptide which comprise contacting the PRO polypeptide with a candidate molecule and monitoring a biological activity mediated by said PRO polypeptide. Preferably, the PRO polypeptide is a native PRO polypeptide.

In a still further embodiment, the invention concerns a composition of matter comprising a PRO polypeptide, or an agonist or antagonist of a PRO polypeptide as herein described, or an anti-PRO antibody, in combination with a carrier. Optionally, the carrier is a pharmaceutically acceptable carrier.

Another embodiment of the present invention is directed to the use of a PRO polypeptide, or an agonist or antagonist thereof as herein before described, or an anti-PRO antibody, for the preparation of a medicament useful in the treatment of a condition which is responsive to the PRO polypeptide, an agonist or antagonist thereof or an anti-PRO antibody.

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Figure 1896: DNA255271, NP_038475.1, 207610_s_at Figure 1946: DNA275286, NP_009205.1, 208002_s_at Figure 1897: PRO50348 Figure 1947: PRO62967 Figure 1898: DNA331498, TANK, 207616_s_at Figure 1948: DNA288217, NP_002101.1, 208018_s_at Figure 1899: PRO86535 Figure 1949: PRO69990 Figure 1900: DNA226337, NP_005683.2, 207622_s_at Figure 1950: DNA227224, NP_060877.1, 208029_s_at Figure 1901: PRO36800 Figure 1951: PRO37687 Figure 1902: DNA227606, NP_001872.2, 207630_s_at Figure 1952A-B: DNA188492, NAB1, 208047_s_at Figure 1903: PRO38069 Figure 1953: PRO22070 Figure 1904: DNA196426, NP_037440.1, 207651_at Figure 1954: DNA330127, NP_006442.2, 208051_s_at Figure 1905: PRO24924 Figure 1955: PRO85387 Figure 1906: DNA328554, NP_038202.1, 207677_s_at Figure 1956A-B: DNA328607, NP_003639.1, Figure 1907: PRO84354 208072_s_at Figure 1908A-B: DNA226405, NP_006525.1, Figure 1957: PRO84390 207700_s_at Figure 1958A-C: DNA331500, NP_003307.2, 208073_x_at Figure 1909: PRO36868 Figure 1910: DNA329064, NP_060301.1, 207735_at Figure 1959: PRO86537 Figure 1911: PRO84724 Figure 1960A-B: DNA328312, NP_110378.1, Figure 1912: DNA329020, NUP62, 207740_s_at 208078_s_at Figure 1913: PRO84695 Figure 1961: PRO84180 Figure 1914: DNA325654, NP_054752.1, 207761_s_at Figure 1962: DNA331501, STK6, 208079_s_at Figure 1915: PRO4348 Figure 1963: PRO58855 Figure 1916A-B: DNA329179, NP_056958.1, Figure 1964: DNA323896, NP_112182.1, 208103_s_at 207785_s_at Figure 1965: PRO80638 Figure 1917: PRO84802 Figure 1966: DNA330129, NP_112495.1, 208119_s_at Figure 1918: DNA227494, NP_002158.1, 207826_s_at Figure 1967: PRO85389 Figure 1919: PRO37957 Figure 1968: DNA325329, NP_004719.1, 208152_s_at Figure 1920A-C: DNA331499, NP_057427.2, Figure 1969: PRO81872 207828_s_at Figure 1970: DNA36717, NP_000581.1, 208193_at Figure 1921: PRO86536 Figure 1971: PRO72 Figure 1922: DNA329182, HPIP, 207838_x_at Figure 1972A-E: DNA330130, HSTITIN, 208195_at Figure 1923: PRO84805 Figure 1973: DNA328611, RASGRP2, 208206_s_at Figure 1924: DNA330123, NP_008984.1, 207840_at Figure 1974: PRO84393 Figure 1925: PRO35080 Figure 1975: DNA328612, NP_000166.2, 208308_s_at Figure 1926: DNA227175, NP_006857.1, 207857_at Figure 1976: PRO84394 Figure 1927: PRO37638 Figure 1977A-D: DNA331502, NP_000050.1, Figure 1928: DNA330124, NP_002981.2, 207861_at 208368_s_at Figure 1929: PRO34107 Figure 1978: PRO86538 Figure 1930: DNA217245, NP_000579.1, 207906_at Figure 1979: DNA324250, NP_536349.1, 208392_x_at Figure 1931: PRO34287 Figure 1980: PRO80934 Figure 1932: DNA218651, NP_003798.1, 207907_at Figure 1981A-B: DNA331503, RAD50, 208393_s_at Figure 1933: PRO34447 Figure 1982: PRO86539 Figure 1934: DNA330125, NP_002729.2, 207957_s_at Figure 1983: DNA327690, NP_004022.1, 208436_s_at Figure 1935: PRO85385 Figure 1984: PRO83673 Figure 1936A-B: DNA226290, NP_036333.1, Figure 1985: DNA103427, NP_005239.1, 208438_s_at 207966_s_at Figure 1986: PRO4755 Figure 1937: PRO36753 Figure 1987A-C: DNA331504, ATM, 208442_s_at Figure 1938: DNA329183, NP_055962.1, 207971_s_at Figure 1988: PRO86540 Figure 1939: PRO84806 Figure 1989A-B: DNA330134, BAZ1B, 208445_s_at Figure 1940A-B: DNA330126, NP_008912.1, Figure 1990: PRO85394 207978_s_at Figure 1991A-C: DNA331505, NP_000642.2, Figure 1941: PRO85386 208488_s_at Figure 1942: DNA329184, CITED2, 207980_s_at Figure 1992: PRO86541 Figure 1943: PRO84807 Figure 1993: DNA330136, NP_002441.1, 208581_x_at Figure 1944A-C: DNA254145, NP_004329.1, Figure 1994: PRO82583 207996_s_at Figure 1995A-C: DNA331506, NP_001448.1, Figure 1945: PRO49260 208614_s_at

Figure 1996: PRO86542 Figure 2051: PRO69498 Figure 1997A-B: DNA330138, PTP4A2, 208617_s_at Figure 2052: DNA329189, NP_009139.1, 208787_at Figure 1998: PRO85397 Figure 2053: PRO4911 Figure 1999A-B: DNA273567, NP_004944.1, Figure 2054: DNA238565, NP_005907.2, 208795_s_at 208624_s_at Figure 2055: PRO39210 Figure 2000: PRO61545 Figure 2056: DNA330145, NP_002788.1, 208799_at Figure 2001A-B: DNA273567, EIF4G1, 208625_s_at Figure 2057: PRO84403 Figure 2002: PRO61545 Figure 2058: DNA331511, HSMPIO, 208805_at Figure 2003: DNA325912, NP_001093.1, 208636_at Figure 2059A-C: DNA331512, 1397486.26, 208806_at Figure 2004: PRO82367 Figure 2060: PRO86547 Figure 2005: DNA325912, ACTN1, 208637_x_at Figure 2061A-B: DNA330147, HSU91543, Figure 2006: PRO82367 208807_s_at Figure 2007: DNA329188, BC012142, 208638_at Figure 2062: PRO85405 Figure 2008: PRO84810 Figure 2063: DNA324531, NP_002120.1, 208808_s_at Figure 2009: DNA324641, NP_005608.1, 208646_at Figure 2064: PRO81185 Figure 2010: PRO10849 Figure 2065: DNA273521, NP_002070.1, 208813_at Figure 2011: DNA271268, NP_009057.1, 208649_s_at Figure 2066: PRO61502 Figure 2012: PRO59579 Figure 2067A-B: DNA330148, AB020636, 208838_at Figure 2013: DNA328617, AF299343, 208653_s_at Figure 2068A-B: DNA330149, HSM801778, Figure 2014: PRO84399 208839_s_at Figure 2015: DNA330139, AK022493, 208657_s_at Figure 2069: PRO82209 Figure 2016: PRO85398 Figure 2070: DNA227874, NP_003320.1, 208864_s_at Figure 2017A-C: DNA151898, TTC3, 208661_s_at Figure 2071: PRO38337 Figure 2018: PRO12135 Figure 2072: DNA328624, BC003562, 208891_at Figure 2019A-C: DNA151898, D84294, 208662_s_at Figure 2073: PRO59076 Figure 2020: PRO12135 Figure 2074: DNA331513, DUSP6, 208892_s_at Figure 2021A-C: DNA331507, D83327, 208663_s_at Figure 2075: PRO84404 Figure 2022: DNA304686, NP_002565.1, 208680_at Figure 2076: DNA331330, BC005047, 208893_s_at Figure 2023: PRO71112 Figure 2077: PRO82215 Figure 2024A-B: DNA328619, BC001188, 208691_at Figure 2078: DNA329221, NP_061984.1, 208894_at Figure 2025: PRO84401 Figure 2079: PRO4555 Figure 2026: DNA287189, NP_002038.1, 208693_s_at Figure 2080A-B: DNA329007, NP_003277.1, Figure 2027: PRO69475 208900_s_at Figure 2028: DNA330140, AF275798, 208696_at Figure 2081: PRO37029 Figure 2029: PRO85399 Figure 2082A-B: DNA329007, TOP1, 208901.s.at Figure 2030A-C: DNA331508, 198777.9, 208707_at Figure 2083: PRO37029 Figure 2031: PRO86543 Figure 2084: DNA327700, BC015130, 208905_at Figure 2032: DNA97298, NP_003899.1, 208726_s_at Figure 2085: PRO83683 Figure 2086: DNA327701, NP_001203.1, 208910_s_at Figure 2033: PRO3645 Figure 2034: DNA330142, BC003564, 208737_at Figure 2087: PRO82667 Figure 2035: PRO85401 Figure 2088: DNA281442, NP_149124.1, 208912_s_at Figure 2036: DNA331509, 1138554.23, 208740_at Figure 2089: PRO66281 Figure 2037: PRO86544 Figure 2090A-B: DNA330151, AB029003, 208914_at Figure 2038: DNA328591, HSP105B, 208744_x_at Figure 2091: DNA325473, NP_006353.2, 208922_s_at Figure 2039: PRO84376 Figure 2092: PRO81996 Figure 2040: DNA287285, NP_005794.1, 208748_s_at Figure 2093: DNA329552, NP_063948.1, 208925_at Figure 2041: PRO69556 Figure 2094: PRO85097 Figure 2042: DNA324217, ATIC, 208758_at Figure 2095: DNA326233, NP_000968.2, 208929_x_at Figure 2043: PRO80908 Figure 2096: PRO82645 Figure 2044: DNA327696, AF228339, 208763_s_at Figure 2097: DNA327702, NP_006490.2, 208934_s_at Figure 2045: PRO83679 Figure 2098: PRO83684 Figure 2046A-B: DNA331510, 1298307.1, 208776_at Figure 2099: DNA330152, NP_001939.1, 208956_x_at Figure 2047: PRO86545 Figure 2100: PRO85406 Figure 2048: DNA287427, NP_002806.1, 208777_s_at Figure 2101: DNA290261, NP_001291.2, 208960_s_at Figure 2049: PRO69684 Figure 2102: PRO70387 Figure 2103A-B: DNA325478, NP_037534.2, Figure 2050: DNA287219, NP_110379.1, 208778_s_at

| 208962_s_at | Figure 2155: PRO84814 |
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| Figure 2104: PRO81999 | Figure 2156A-B: DNA330161, NP_085059.1, |
| Figure 2105: DNA327661, IFI16, 208965_s_at | 209081_s_at |
| Figure 2106: PRO83652 | Figure 2157: PRO85413 |
| Figure 2107A-B: DNA270277, AF208043, | Figure 2158: DNA330162, NP_057093.1, 209091_s_at |
| 208966_x_at | Figure 2159: PRO85414 |
| Figure 2108: PRO58665 | Figure 2160: DNA330163, NP_060308.1, 209104_s_at |
| Figure 2109: DNA326343, KPNB1, 208974_x_at | Figure 2161: PRO85415 |
| Figure 2110: PRO82739 | Figure 2162: DNA330164, NP_004752.1, 209110_s_at |
| Figure 2111A-B: DNA330153, HUMIMP90A, | Figure 2163: PRO85416 |
| 208975_s_at | Figure 2164: DNA327709, NP_000509.1, 209116_x_at |
| Figure 2112: PRO82739 | Figure 2165: PRO83690 |
| Figure 2113: DNA328629, NP_006079.1, 208977_x_at | Figure 2166: DNA288254, NP_006000.2, 209118_s_at |
| Figure 2114: PRO84407 | Figure 2167: PRO69536 |
| Figure 2115: DNA330154, HUMPECAM27, | Figure 2168: DNA325163, NP_001113.1, 209122_at |
| 208981_at | Figure 2169: PRO81730 |
| Figure 2116: DNA330155, 7692317.2, 208982_at | Figure 2170: DNA330165, BC015833, 209138_x_at |
| Figure 2117: PRO85407 | Figure 2171: PRO85417 |
| Figure 2118: DNA330156, NP_003749.1, 208985_s_at | Figure 2172: DNA327713, BC010653, 209146_at |
| Figure 2119: PRO85408 | Figure 2173: PRO37975 |
| Figure 2120: DNA331514, STAT3, 208992_s_at | Figure 2174: DNA325285, AKR1C3, 209160_at |
| Figure 2121: PRO86548 | Figure 2175: PRO81832 |
| Figure 2122: DNA227552, NP_003346.2, 208997_s_at | Figure 2176: DNA330166, BC001588, 209161_at |
| Figure 2123: PRO38015 | Figure 2177: PRO85418 |
| Figure 2124: DNA227552, UCP2, 208998_at | Figure 2178: DNA271722, NP_004688.1, 209162_s_at |
| Figure 2125: PRO38015 | Figure 2179: PRO60006 |
| Figure 2126: DNA328630, NP_036293.1, 209004_s_at | Figure 2180: DNA330167, CAB43224.1, 209177_at |
| Figure 2127: PRO84408 | Figure 2181: PRO85419 |
| Figure 2128: DNA331515, FBXL5, 209005.at Figure 2129: PRO86549 | Figure 2182A-B: DNA328642, AF073310, 209184_s_at |
| Figure 2130: DNA328631, AK027318, 209006_s_at | Figure 2183: PRO84418 |
| Figure 2131: PRO84409 | Figure 2184: DNA331331, AF161416, 209185_s_at |
| Figure 2132: DNA331516, DNAJB6, 209015_s_at | Figure 2185A-B: DNA328643, HUMHK1A, |
| Figure 2133: PRO83680 | 209186_at |
| Figure 2134: DNA328633, NP_004784.2, 209017_s_at | Figure 2186: PRO84419 |
| Figure 2135: PRO84411 | Figure 2187: DNA189700, NP_005243.1, 209189_at |
| Figure 2136: DNA330158, NP_057554.4, 209020_at | Figure 2188: PRO25619 |
| Figure 2137: PRO85410 | Figure 2189: DNA324766, NP_005443.2, 209196_at |
| Figure 2138: DNA327851, NP_006363.2, 209024_s_at | Figure 2190: PRO81387 |
| Figure 2139: PRO83795 | Figure 2191: DNA226176, NP_003458.1, 209201_x_at |
| Figure 2140: DNA328635, BC020946, 209026_x_at | Figure 2192: PRO36639 |
| Figure 2141: PRO84413 | Figure 2193: DNA326267, NP_004861.1, 209208_at |
| Figure 2142: DNA331517, NP_004150.1, 209040_s_at | Figure 2194: PRO82674 |
| Figure 2143: PRO69506 | Figure 2195: DNA326891, NP_001748.1, 209213_at |
| Figure 2144A-C: DNA328637, HSA7042, 209052_s_at | Figure 2196: PRO83212 |
| Figure 2145: PRO81109 | Figure 2197: DNA227483, NP_003120.1, 209218_at |
| Figure 2146A-B: DNA331518, AF330040, | Figure 2198: PRO37946 |
| 209053_s_at | Figure 2199: DNA330168, NP_006322.1, 209233_at |
| Figure 2147: PRO86550 | Figure 2200: PRO85420 |
| Figure 2148A-B: DNA226405, NCOA3, 209060_x_at | Figure 2201: DNA328649, NP_116093.1, 209251_x_at |
| Figure 2149: PRO36868 | Figure 2202: PRO84424 |
| Figure 2150: DNA330159, HSM801885, 209064_x_at | Figure 2203: DNA255255, NP_071437.1, 209267_s_at |
| Figure 2151: PRO85411 | Figure 2204: PRO50332 |
| Figure 2152: DNA330160, NP_006285.1, 209066_x_at | Figure 2205A-B: DNA188492, AF045451, 209272_at |
| Figure 2153: PRO85412 Figure 2154: DNA329194, NP_112740.1, 209068_at | Figure 2206: PRO22070 Figure 2207A-B: DNA226827, NP_001673.1, |
| FIRUIC 4134. DINA347174, INF_114/40.1, 209008_8[| rigule 220/M-D. DIMA22002/, INF_0010/3.1, |

| Figure 2209: PRO37290 Figure 2209: PRO37290 Figure 2209: DNA328601, GADD45B, 209304_x.at Figure 2210: PRO84384 Figure 2211: DNA328651, AF087853, 209305_s.at Figure 2212: PRO84389 Figure 2212: PRO52889 Figure 2213: DNA151780, NP_006611.1, 209314_s.at Figure 2214: PRO12057 Figure 2215: DNA230169, NP_006709_1, 209318_x.at Figure 2216: PRO52736 Figure 2217: DNA275106, HSU76248, 209339_at Figure 2218: PRO52821 Figure 2219: PNA287506, HSU76248, 209339_at Figure 2219: PNA287506, HSU76248, 209339_at Figure 22219: PRO58042 Figure 2222: PRO58042 Figure 2222: PRO58042 Figure 2222: PRO58043 Figure 2223: DNA330170, AF109161, 209357_at Figure 2222: PRO58432 Figure 2223: DNA330171, CAA34971.1, 209374_s.at Figure 2223: PRO58542 Figure 2223: PRO58542 Figure 2239: DNA330173, HUMAUTOTAX, 20939_at Figure 2239: DNA330173, HUMAUTOTAX, 20939_at Figure 2239: DNA330173, HUMAUTOTAX, 20939_at Figure 2239: DNA330175, NP_006836.1, 20940_s.at Figure 2239: DNA330177, BC000529, 209377_s.at Figure 2239: PRO59562 Figure 2239: DNA330178, BTMA2668, AP064943, 209585_s.at Figure 2239: DNA330178, BTMA2668, AP064948_at Figure 2239: DNA330179, BC000748, AP064948_at Figure 2239: DNA330179, BC000748, AP064948_at Figure 2240: PRO65426 Figure 2241: DNA27569, BA2, 209408_at Figure 22429: PRO65426 Figure 22429: PRO65426 Figure 22429: DNA330179, BC00748, 209408_at Figure 22429: PRO65436 Figure 22429: PRO65436 F | | |
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| Figure 2209: DNA328601, GADD45B, 209304.x.t Figure 2209: PRO84344 Figure 2211: DNA328651, AF087853, 209305.s.at Figure 2211: DNA328651, AF087853, 209305.s.at Figure 2212: DNA330180, NP.006101.1, 209314.s.at Figure 2216: DNA320169, NP.006709.1, 209318.x.at Figure 2216: PRO62736 Figure 2217: DNA275106, HSU76248, 209339.at Figure 2216: DNA275060, NP.003281.1, 209344.at Figure 2220: DNA269630, NP.003281.1, 209344.at Figure 2220: DNA269630, NP.003281.1, 209344.at Figure 2221: DNA2530170, AF109161, 209357.at Figure 2222: DNA330170, AF109161, 209357.at Figure 2223: DNA330170, AF109161, 209357.at Figure 2223: DNA330170, AF109161, 209357.at Figure 2223: DNA330171, CAA34971.1, 209374.s.at Figure 2224: PRO84807 Figure 2225: DNA330173, HUMAUTOTAX, 209392.at Figure 2232: DNA330173, HUMAUTOTAX, 209392.at Figure 2232: DNA330174, AK027512, 209404.s.at Figure 2234: DNA330173, HUMAUTOTAX, 209392.at Figure 2235: DNA330174, AK027512, 209404.s.at Figure 2236: DNA330175, NP.006836.1, 209408.at Figure 2236: DNA328666, AF084943, 209585.s.at Figure 2236: DNA330176, AAB61703.1, 209417.s.at Figure 2236: DNA330176, AB61703.1, 209417.s.at Figure 2246: PRO85426 Figure 2237: DNA330176, AAB61703.1, 209417.s.at Figure 2246: PRO85426 Figure 2237: DNA330178, NP.006836.1, 209408.s.at Figure 2249: DNA25767, NP.000534.1, 209408.s.at Figure 2239: DNA330176, AAB61703.1, 209417.s.at Figure 2249: PRO3629 PRO59568 Figure 2239: DNA330178, NP.000534.1, 209408.s.at Figure 2239: DNA330178, NP.000534.1, 209404.s.at Figure 2236: DNA330178, NP.0074279.2, 209606.at Figure 2237: DNA330183, NP.004279.2, 209606.at Figure 2239: DNA330178, NP.006836.1, 209408.s.at Figure 2239: DNA330178, NP.006838.3, 209451.at Figure 2239: DNA330183, NP.004279. | 209281_s_at | Figure 2261: DNA274027, RAB27A, 209514_s_at |
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| Figure 2319: PRO85439 | Figure 2373: DNA154921, DNA154921, 209967_s_at |
| Figure 2320A-C: DNA254412, AF008915, 209717_at | Figure 2374: DNA327736, BC002704, 209969_s_at |
| Figure 2321: PRO49522 | Figure 2375: PRO83711 |
| Figure 2322A-B: DNA330192, 234780.1, 209733.at | Figure 2376: DNA324895, JTV1, 209971_x_at |
| Figure 2323: PRO85440 | Figure 2377: PRO81501 |
| Figure 2324: DNA330193, BC015929, 209750_at | Figure 2378: DNA226658, NP_003736.1, 209999_x_at |
| Figure 2325: DNA330194, HSU09087, 209754_s_at | Figure 2379: PRO37121 |
| Figure 2326: PRO85442 | Figure 2380: DNA226658, SSI-1, 210001_s_at |
| Figure 2327: DNA275195, NP_001025.1, 209773_s_at | Figure 2381: PRO37121 |
| Figure 2328: PRO62893 | Figure 2382: DNA330200, NP_056222.1, 210006_at |
| Figure 2329: DNA329205, NP_001343.1, 209782.s_at | Figure 2383: PRO85448 |
| Figure 2330: PRO84821 | Figure 2384: DNA269534, NP_002155.1, 210029_at |
| Figure 2331: DNA226436, NP_001772.1, 209795_at | Figure 2385: PRO57950 |
| Figure 2332: PRO36899 | Figure 2386: DNA326054, NP_002159.1, 210046_s_at |
| Figure 2333: DNA327731, NP_003302.1, 209803_s_at | Figure 2387: PRO82489 |
| Figure 2334: PRO83707 | Figure 2388: DNA326809, NP_036244.2, 210052_s_at |
| Figure 2335A-B: DNA271368, HUMKIAAI, | Figure 2389: PRO83142 |
| 209804_at | Figure 2390: DNA150551, AAB97010.1, 210054_at |
| Figure 2336: PRO59668 | Figure 2391: PRO12778 |
| Figure 2337: DNA329206, AF151103, 209813_x_at | Figure 2392: DNA274960, SFRS5, 210077_s_at |
| Figure 2338: PRO84822 | Figure 2393: PRO62694 |
| Figure 2339A-C: DNA331520, 1096711.1, 209815_at | Figure 2394: DNA324922, BC018962, 210095_s_at |
| Figure 2340: PRO86552 | Figure 2395: PRO119 |
| Figure 2341: DNA327732, UMPK, 209825_s_at | Figure 2396A-B: DNA328685, NP_127497.1, |
| Figure 2342: PRO61801 | 210113_s_at |
| Figure 2343: DNA328676, IL16, 209827.s_at | Figure 2397: PRO34751 |
| Figure 2344: PRO84448 | Figure 2398: DNA330201, NP_003774.1, 210121_at |
| Figure 2345A-B: DNA196499, AB002384, 209829_at | Figure 2399: PRO50625 |
| Figure 2346: PRO24988 | Figure 2400: DNA330202, NP_005400.1, 210163_at |
| Figure 2347: DNA330197, NP_112190.1, 209832_s_at | Figure 2401: PRO19838 |
| Figure 2348: PRO85445 | Figure 2402: DNA287620, NP_004122.1, 210164_at |
| Figure 2349: DNA328677, AF060511, 209836_x_at | Figure 2403: PRO2081 Figure 2404: DNA270196, HUMZFM1B, 210172_at |
| Figure 2350: PRO84449 Figure 2351: DNA270180, NP_478123.1, 209849_s_at | Figure 2405: PRO58584 |
| Figure 2352: PRO58569 | Figure 2406: DNA330203, NP_003755.1, 210190_at |
| Figure 2353: DNA331521, BC018951, 209868_s_at | Figure 2407: PRO85449 |
| Figure 2354: PRO58719 | Figure 2408: DNA331335, AF070576, 210201_x_at |
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| Figure 2357: DNA330198, AB014719, 209871_s_at | Figure 2412: PRO61690 |
| Figure 2358: PRO85446 | Figure 2413: DNA328467, AF056322, 210218_s_at |
| Figure 2359: DNA324184, NP_065726.1, 209891_at | Figure 2414: PRO84293 |
| Figure 2360: PRO80882 | Figure 2415: DNA217253, NP_000749.1, 210229_s_at |
| Figure 2361: DNA328258, HSM802616, 209900_s_at | Figure 2416: PRO34295 |
| Figure 2362: PRO84151 | Figure 2417: DNA331084, BC008487, 210254_at |
| Figure 2363: DNA330152, DUT, 209932_s_at | Figure 2418: PRO81984 |
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| Figure 2365: DNA150133, AAD01646.1, 209933_s_at | 210281_s_at |
| Figure 2366: PRO12219 | Figure 2420: PRO58410 |
| Figure 2367: DNA329208, CFLAR, 209939_x_at | Figure 2421: DNA330206, NP_005801.2, 210288_at |
| Figure 2368: PRO84823 | Figure 2422: PRO85450 |
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| Figure 2424: PRO85252 | Figure 2476: DNA330215, DKFZp762A227Homo, |
| Figure 2425: DNA218653, NP_003799.1, 210314_x_at | 210692_s_at |
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| Figure 2427: DNA326239, NP_006752.1, 210317_s_at | Figure 2478: PRO86559 |
| Figure 2428: PRO39530 | Figure 2479: DNA237817, NP_001307.1, 210766_s_at |
| Figure 2429: DNA329213, NP_219491.1, 210321_at | Figure 2480: PRO38923 |
| Figure 2430: PRO2313 | Figure 2481A-B: DNA330216, NP_006445.1, |
| Figure 2431A-B: DNA329214, NP_001087.1, | 210778_s_at |
| 210337_s_at | Figure 2482: PRO85457 |
| Figure 2432: PRO84826 | Figure 2483: DNA226881, FLI1, 210786_s_at |
| Figure 2433: DNA225528, NP_000610.1, 210354_at | Figure 2484: PRO37344 |
| Figure 2434: PRO35991 | Figure 2485: DNA255402, NP_055288.1, 210802_s_at |
| Figure 2435: DNA196621, HUMLY9, 210370_s_at | Figure 2486: PRO50469 |
| Figure 2436: DNA330207, BC001131, 210387_at | Figure 2487: DNA330027, SSBP2, 210829_s_at |
| Figure 2437: PRO85451 | Figure 2488: PRO85312 |
| Figure 2438: DNA226229, NP_002432.1, 210410_s_at | Figure 2489: DNA329219, BC000385, 210844_x_at |
| Figure 2439: PRO36692 | Figure 2490: PRO81278 |
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| Figure 2444: DNA226394, NP_002552.1, 210448_s_at | Figure 2495: DNA331531, PFDN5, 210908_s_at |
| Figure 2445: PRO36857 | Figure 2496: PRO86560 |
| Figure 2446: DNA331524, BC003388, 210458_s_at Figure 2447: PRO86554 | Figure 2497: DNA330217, AF043183, 210915_x_at |
| Figure 2448: DNA331525, BC002448, 210461_s_at | Figure 2498: PRO85458 Figure 2499: DNA274326, NP_003079.1, 210933_s_at |
| Figure 2449: PRO86555 | Figure 2500: PRO62244 |
| Figure 2450: DNA329216, AF022375, 210512_s_at | Figure 2501: DNA329317, NP_057353.1, 210948_s_at |
| Figure 2451: PRO84827 | Figure 2502: PRO81157 |
| Figure 2452: DNA227633, NP_001156.1, 210538_s_at | Figure 2503: DNA331532, AF125393, 210951_x_at |
| Figure 2453: PRO38096 | Figure 2504: PRO86561 |
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| Figure 2456: DNA331526, BC014563, 210559_s_at | Figure 2506: DNA273236, NP_004306.1, 210980_s_at |
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| Figure 2409: PRO83455 Figure 2470: DNA331529, LAIR1, 210644_s_at | Figure 2519: PRO83682 Figure 2520: DNA 288254, TUBA 3, 211058, v. et |
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| Figure 2527: PRO60769 | Figure 2580: DNA226342, PTEN, 211711_s_at |
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| Figure 2540: PRO84464 | Figure 2595: PRO85462 Figure 2596A-B: DNA329226, BC006181, |
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| Figure 3164A-B: DNA254376, KIAA0963, | Figure 3217A-B: DNA330331, BAA86451.1, |
| 215760_s_at | 216873_s_at |
| Figure 3165: PRO49486 | Figure 3218: PRO85554 |
| Figure 3166A-B: DNA328805, BAA86482.1, | Figure 3219: DNA188275, NP_002181.1, 216876_s_at |
| 215785_s_at | Figure 3220: PRO21800 |
| Figure 3167: PRO84547 | Figure 3221A-B: DNA150987, NP_006051.1, |
| Figure 3168: DNA331574, HUMTCGCH, 215806_x_at | 216901_s_at |
| Figure 3169: DNA330322, 234025.21, 215855_s_at | Figure 3222: PRO12163 |
| Figure 3170: PRO85546 | Figure 3223: DNA329267, HUMTCRGAAC, |
| Figure 3171: DNA330323, 335053.1, 215908_at | 216920_s_at |
| Figure 3172: PRO85547 | Figure 3224A-C: DNA328811, HUMINSP3R1, |
| Figure 3173: DNA330324, HHEX, 215933_s_at | 216944_s_at |
| Figure 3174: PRO58034 | Figure 3225: PRO84551 |
| Figure 3175: DNA331575, AF223408, 215942.s.at | Figure 3226A-B: DNA151027, AAA80979.1, |
| Figure 3176: PRO86587 | 216952_s_at |
| Figure 3177: DNA330325, NP_055057.1, 215948_x_at | Figure 3227: PRO12843 |
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| | Figure 3228A-E: DNA269650, PLEC1, 2169/1.S.at |
| | Figure 3228A-E: DNA269650, PLEC1, 216971_s_at Figure 3229A-B: PRO58061 |
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| Figure 3257: DNA329539, HLA-DMA, 217478_s_at | Figure 3311: PRO49871 |
| Figure 3258: PRO85089 | Figure 3312: DNA330342, NP_067021.1, 217844_at |
| Figure 3259: DNA331589, 243999.2, 217502_at | Figure 3313: PRO85564 |
| Figure 3260: PRO86596 | Figure 3314: DNA329272, NP_055181.1, 217850_at |
| Figure 3261: DNA329271, 406848.1, 217591_at | Figure 3315: PRO84868 |
| Figure 3262: PRO84867 | Figure 3316A-B: DNA330343, AF403012, |
| Figure 3263: DNA330337, 1447003.1, 217616_at | 217857_s_at |
| Figure 3264: PRO85559 | Figure 3317: DNA330344, NP_057392.1, 217870_s_at |
| Figure 3265: DNA331590, 368556.1, 217655_at | Figure 3318: PRO85565 |
| Figure 3266: PRO86597 | Figure 3319: DNA326937, NP_002406.1, 217871_s_at |
| Figure 3267: DNA330339, HSA012375, 217672_x_at | Figure 3320: PRO83255 |
| Figure 3268: DNA323856, HSM800628, 217725_x_at | Figure 3321: DNA330345, NP_055130.1, 217906_at |
| Figure 3269: PRO80599 | Figure 3322: PRO85566 |
| Figure 3270: DNA326523, NP_001121.2, 217729_s_at Figure 3271: PRO71126 | Figure 3323: DNA330346, NP_054880.2, 217907_at |
| Figure 3271: PRO71120 Figure 3272: DNA325832, NP_068839.1, 217731_s_at | Figure 3324: PRO85567 Figure 3325: DNA325780, NP_060371.1, 217914_at |
| Figure 3273: PRO1869 | Figure 3326: PRO82250 |
| Figure 3274A-B: DNA327847, 142131.14, 217738_at | Figure 3327: DNA327853, NP_054769.1, 217919_s_at |
| Figure 3275: PRO2834 | Figure 3328: PRO82223 |
| Figure 3276: DNA88541, NP_005737.1, 217739_s_at | Figure 3329: DNA330347, 255559.4, 217922_at |
| Figure 3277: PRO2834 | Figure 3330: PRO85568 |
| Figure 3278: DNA327935, NP_079422.1, 217745_s_at | Figure 3331: DNA330348, NP_079150.1, 217929_s_at |
| Figure 3279: PRO83866 | Figure 3332: PRO85569 |
| Figure 3280: DNA327849, NP_057269.1, 217755_at | Figure 3333: DNA330349, BC022093, 217931_at |
| Figure 3281: PRO83794 | Figure 3334: DNA287241, NP_056991.1, 217933_s_at |
| Figure 3282A-B: DNA274131, AF183421, | Figure 3335: PRO69516 |
| 217762_s_at | Figure 3336A-B: DNA225648, NP_061165.1, |
| Figure 3283: PRO62067 | 217941_s_at |
| Figure 3284: DNA330340, NP_006859.1, 217763_s_at | Figure 3337: PRO36111 |
| Figure 3285: PRO85562 | Figure 3338: DNA326730, NP_057037.1, 217950_at |
| Figure 3286A-B: DNA274131, DNA274131, | Figure 3339: PRO83072 |
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| Figure 3287: PRO62067 | Figure 3341: PRO84869 |
| Figure 3288: DNA325821, NP_057016.1, 217769_s_at | Figure 3342: DNA328829, NP_057230.1, 217959_s_at |
| Figure 3289: PRO82287 | Figure 3343: PRO84566 |
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| Figure 3293: PRO37821 | Figure 3347: PRO84870 |
| Figure 3294: DNA328819, NP_057145.1, 217783_s_at | Figure 3348: DNA325496, NP_037397.2, 217969_at |
| Figure 3295: PRO84557 | Figure 3349: PRO82013 |
| Figure 3296: DNA325873, SKB1, 217786_at | Figure 3350: DNA327855, NP_057387.1, 217975_at |
| Figure 3297: PRO82331 | Figure 3351: PRO83367 |
| Figure 3298: DNA331591, NP_055241.1, 217792_at | Figure 3352: DNA227218, NP_003721.2, 217983_s_at |
| Figure 3299: PRO69560 | Figure 3353: PRO37681 |
| Figure 3300: DNA328303, NP_056525.1, 217807_s_at | Figure 3354: DNA227218, RNASE6PL, 217984_at |
| Figure 3301: PRO84173 | Figure 3355: PRO37681 |
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Figure 3357: PRO37701 Figure 3413: DNA304470, PRO2577, 218172_s_at Figure 3358A-B: DNA227238, BAZ1A, 217986_s_at Figure 3414: PRO2577 Figure 3359: PRO37701 Figure 3415: DNA330359, NP_065145.1, 218178_s_at Figure 3360: DNA328831, NP_057329.1, 217989_at Figure 3416: PRO85575 Figure 3361: PRO233 Figure 3417: DNA304495, NP_057156.1, 218193_s_at Figure 3362: DNA328832, NP_067022.1, 217995_at Figure 3418: PRO793 Figure 3363: PRO84568 Figure 3419A-C: DNA330360, NP_078789.1, Figure 3364: DNA328833, BC018929, 217996_at 218204_s_at Figure 3365: PRO84569 Figure 3420: PRO85576 Figure 3366: DNA328834, AF220656, 217997_at Figure 3421: DNA327858, NP_036473.1, 218238_at Figure 3367: DNA287364, NP_031376.1, 218000_s_at Figure 3422: PRO83800 Figure 3368: PRO69625 Figure 3423: DNA327858, CRFG, 218239_s_at Figure 3369: DNA273008, NP_003972.1, 218009_s_at Figure 3424: PRO83800 Figure 3370: PRO61079 Figure 3425A-B: DNA330361, CKAP2, 218252_at Figure 3371: DNA330350, NP_006108.1, 218025_s_at Figure 3426: PRO85577 Figure 3372: PRO85570 Figure 3427: DNA328850, NP_057187.1, 218254_s_at Figure 3373: DNA328836, NP_054894.1, 218027_at Figure 3428: PRO84581 Figure 3374: PRO84572 Figure 3429: DNA331594, MRPL24, 218270_at Figure 3375: DNA329275, AF070673, 218032_at Figure 3430: PRO11652 Figure 3376: PRO12342 Figure 3431: DNA273230, NP_060914.1, 218273_s_at Figure 3377: DNA331592, ANKT, 218039_at Figure 3432: PRO61257 Figure 3378: PRO82424 Figure 3433: DNA324444, NP_006333.1, 218308_at Figure 3379: DNA328838, NP_054797.2, 218049_s_at Figure 3434: PRO81108 Figure 3380: PRO70319 Figure 3435: DNA330363, NP_060252.1, 218331_s_at Figure 3381: DNA330352, NP_075059.1, 218051_s_at Figure 3436: PRO85578 Figure 3437: DNA329281, NP_036526.2, 218336_at Figure 3382: PRO85571 Figure 3383: DNA329276, NP_077001.1, 218069_at Figure 3438: PRO84874 Figure 3384: PRO12104 Figure 3439A-B: DNA330364, NP_004417.1, Figure 3385: DNA328841, NP_060557.2, 218073_s_at 218338_at Figure 3386: PRO84575 Figure 3440: PRO85579 Figure 3387: DNA329277, NP_054883.3, 218084_x_at Figure 3441: DNA272918, NP_055269.1, 218346_s_at Figure 3388: PRO6241 Figure 3442: PRO61003 Figure 3389: DNA330353, BC020796, 218085_at Figure 3443: DNA327862, NP_060445.1, 218349_s_at Figure 3390: PRO69464 Figure 3444: PRO83803 Figure 3391: DNA329278, NP_004495.1, 218092_s_at Figure 3445: DNA328854, NP_056979.1, 218350_s_at Figure 3392: PRO84871 Figure 3446: PRO84585 Figure 3393: DNA227313, NP_060945.1, 218095_s_at Figure 3447A-B: DNA273415, KIF4A, 218355_at Figure 3394: PRO37776 Figure 3448: PRO61414 Figure 3395: DNA331593, MRPL4, 218105_s_at Figure 3449: DNA324890, NP_037525.1, 218356_at Figure 3396: PRO86598 Figure 3450: PRO81496 Figure 3397: DNA326596, NP_060624.1, 218115.at Figure 3451: DNA330365, NP_036591.1, 218357_s_at Figure 3398: PRO82954 Figure 3452: PRO85580 Figure 3399: DNA330355, NP_055063.1, 218117_at Figure 3453A-B: DNA331595, NP_073602.2, Figure 3400: PRO83289 218376_s_at Figure 3401: DNA330356, NP_006318.1, 218118_s_at Figure 3454: PRO86599 Figure 3402: PRO85572 Figure 3455: DNA330367, NP_057174.1, 218379_at Figure 3403: DNA330357, NP_078786.2, 218130_at Figure 3456: PRO85582 Figure 3404: PRO85573 Figure 3457: DNA328856, NP_068376.1, 218380_at Figure 3405: DNA227155, NP_057654.1, 218135_at Figure 3458: PRO84586 Figure 3406: PRO37618 Figure 3459: DNA227248, NP_006287.1, 218397_at Figure 3407: DNA254496, NP_060076.1, 218149_s_at Figure 3460: PRO37711 Figure 3408: PRO49604 Figure 3461A-B: DNA287192, NP_006178.1, Figure 3409: DNA330358, NP_079012.1, 218154_at 218400_at Figure 3410: PRO85574 Figure 3462: PRO69478 Figure 3411: DNA254739, NP_068766.1, 218156_s_at Figure 3463: DNA329912, TTC4, 218442_at Figure 3412: PRO49837 Figure 3464: PRO85227

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Figure 3683: DNA271811, NP_036514.1, 219421_at Figure 3734: PRO37718 Figure 3684: PRO60092 Figure 3735: DNA328919, NP_078987.1, 219777_at Figure 3685: DNA329014, NP_005746.2, 219424_at Figure 3736: PRO84637 Figure 3686: PRO9998 Figure 3737A-B: DNA331602, NP_060568.3, Figure 3687: DNA328901, FLJ20533, 219449_s_at 219787_s_at Figure 3738: PRO86604 Figure 3688: PRO84622 Figure 3689: DNA328902, NP_071750.1, 219452_at Figure 3739: DNA255822, NP_036346.1, 219797_at Figure 3690: PRO84623 Figure 3740: PRO50877 Figure 3691: DNA328367, RIN3, 219456_s_at Figure 3741: DNA227305, NP_064564.1, 219806_s_at Figure 3692: PRO84226 Figure 3742: PRO37768 Figure 3693: DNA331598, AK026092, 219457_s_at Figure 3743: DNA329303, NP_054737.1, 219819_s_at Figure 3694: PRO86601 Figure 3744: PRO84892 Figure 3695: DNA327890, NP_079021.1, 219493_at Figure 3745: DNA287295, NP_078784.1, 219836_at Figure 3696: PRO83826 Figure 3746: PRO69564 Figure 3747: DNA287234, NP_114174.1, 219862_s_at Figure 3697A-B: DNA227179, NP_059120.1, 219505_at Figure 3748: PRO69513 Figure 3698: PRO37642 Figure 3749: DNA287221, NP_057407.1, 219863_at Figure 3699A-C: DNA331599, BCL11B, 219528_s_at Figure 3750: PRO69500 Figure 3700: PRO86602 Figure 3751: DNA330419, NP_038469.1, 219864_s_at Figure 3701: DNA329300, GEMIN6, 219539_at Figure 3752: PRO85624 Figure 3702: PRO84889 Figure 3753: DNA255255, LOC64116, 219869_s_at Figure 3703: DNA328908, 7691567.2, 219540_at Figure 3754: PRO50332 Figure 3704: PRO84629 Figure 3755: DNA330420, NP_078890.1, 219871_at Figure 3705: DNA256737, NP_060276.1, 219541_at Figure 3756: PRO85625 Figure 3757: DNA256325, NP_005470.1, 219889_at Figure 3706: PRO51671 Figure 3707: DNA330410, NP_060925.1, 219555_s_at Figure 3758: PRO51367 Figure 3708: PRO85618 Figure 3759: DNA330421, NP_057438.2, 219911_s_at Figure 3709A-B: DNA331600, NP_061985.1, Figure 3760: PRO85626 219577_s_at Figure 3761A-B: DNA330422, NP_057736.2, Figure 3710: PRO86603 219913_s_at Figure 3711: DNA325053, NP_060230.2, 219588_s_at Figure 3762: PRO85627 Figure 3712: PRO81637 Figure 3763: DNA227787, NP_060606.1, 219918.s_at Figure 3713: DNA330412, NP_057617.1, 219594_at Figure 3764: PRO38250 Figure 3714: PRO23600 Figure 3765: DNA330423, NP_037466.2, 219920_s_at Figure 3715: DNA331601, NP_071915.1, 219628_at Figure 3766: PRO85628 Figure 3716: PRO85620 Figure 3767A-B: DNA330424, LTBP3, 219922_s_at Figure 3717: DNA330414, NP_057615.1, 219657.s.at Figure 3768: PRO85629 Figure 3718: PRO81138 Figure 3769: DNA328924, NP_057150.2, 219933_at Figure 3719A-B: DNA274044, HSM801565, Figure 3770: PRO84641 219671_at Figure 3771: DNA218280, NP_068570.1, 219971_at Figure 3720: PRO61987 Figure 3772: PRO34332 Figure 3721: DNA293243, RCP, 219681_s_at Figure 3773: DNA325979, NP_060924.4, 219978_s_at Figure 3722: PRO70699 Figure 3774: PRO82424 Figure 3775: DNA330425, NP_078956.1, 219990_at Figure 3723: DNA255161, NP_071430.1, 219684_at Figure 3724: PRO50241 Figure 3776: PRO85630 Figure 3725: DNA287206, NP_060124.1, 219691_at Figure 3777A-B: DNA330426, SGKL, 220038_at Figure 3726: PRO69488 Figure 3778: PRO85631 Figure 3727A-B: DNA330297, NP_065138.2, Figure 3779: DNA328926, NP_064703.1, 220046_s_at 219700_at Figure 3780: PRO84643 Figure 3728: PRO85524 Figure 3781A-B: DNA218680, NP_071731.1, Figure 3729: DNA330416, TDP1, 219715_at 220048_at Figure 3730: PRO85622 Figure 3782: PRO21724 Figure 3731: DNA330417, NP_085144.1, 219716_at Figure 3783: DNA330427, NP_036593.1, 220052_s_at Figure 3732: PRO21341 Figure 3784: PRO85632 Figure 3733A-B: DNA227255, NP_036579.1, Figure 3785: DNA330428, NP_060385.1, 220060_s_at 219753_at Figure 3786: PRO85633

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| Figure 3789: DNA256091, NP_071385.1, 220094_s_at | Figure 3843: PRO85643 |
| Figure 3790: PRO51141 | Figure 3844: DNA330443, NP_061086.1, 220702_at |
| Figure 3791: DNA330430, NP_078945.1, 220112_at | Figure 3845: PRO85644 |
| Figure 3792: PRO85634 | Figure 3846: DNA288247, NP_478059.1, 220892_s_at |
| Figure 3793: DNA330431, NP_055198.1, 220118_at | Figure 3847: PRO70011 |
| Figure 3794: PRO85635 | Figure 3848: DNA327916, NP_079466.1, 220940_at |
| Figure 3795: DNA227302, NP_037401.1, 220132_s_at | Figure 3849: PRO83851 |
| Figure 3796: PRO37765 | Figure 3850: DNA327953, NP_055182.2, 220942_x_at |
| Figure 3797: DNA330432, NP_079219.1, 220169_at | Figure 3851: PRO83878 |
| Figure 3798: PRO85636 | Figure 3852: DNA327917, NP_112240.1, 220966_x_at |
| Figure 3799: DNA331603, TMPRSS3, 220177_s_at | Figure 3853: PRO83852 |
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| Figure 3801: DNA256291, NP_079182.1, 220232_at | Figure 3855: PRO23253 |
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| Figure 3803: DNA330434, NP_060842.1, 220235_s_at Figure 3804: PRO85637 | 220992_s_at |
| Figure 3805: DNA330435, NP_060179.1, 220306_at | Figure 3857: PRO49623 |
| Figure 3806: PRO85638 | Figure 3858: DNA330444, NP_110405.1, 220999_s_at Figure 3859: PRO85645 |
| Figure 3807: DNA330436, NP_037394.1, 220319_s_at | |
| Figure 3808: PRO85639 | Figure 3860: DNA324246, NP_112188.1, 221004_s_at Figure 3861: PRO80930 |
| Figure 3809: DNA327904, NP_071419.2, 220330_s_at | Figure 3862: DNA330445, NP_112174.1, 221012_s_at |
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| Figure 3816: PRO50795 | Figure 3869: PRO85648 |
| Figure 3817: DNA327214, NP_078991.2, 220495_s_at | Figure 3870: DNA226227, NP_060872.1, 221111_at |
| Figure 3818: PRO83483 | Figure 3871: PRO36690 |
| Figure 3819: DNA324252, NP_060444.1, 220521_s_at | Figure 3872: DNA227267, NP_061130.1, 221123_x_at |
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| Figure 3825: DNA255798, NP_079265.1, 220576_at | Figure 3878: DNA324408, NP_060493.2, 221203_s_at |
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| Figure 3827: DNA330440, NP_079098.1, 220591_s_at | Figure 3880A-B: DNA330448, NP_059111.1, |
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| Figure 3829: DNA255734, NP_057607.1, 220646_s_at | Figure 3881: PRO85649 |
| Figure 3830: PRO50791 | Figure 3882: DNA331605, CISH, 221223_x_at |
| Figure 3831A-B: DNA327908, MCM10, 220651_s_at | Figure 3883: PRO86458 |
| Figure 3832: PRO83843 | Figure 3884: DNA330450, AK025947, 221235_s_at |
| Figure 3833: DNA329306, NP_079149.2, 220655_at Figure 3834: PRO84895 | Figure 3885: PRO85651 |
| Figure 3835A-B: DNA327909, ARNTL2, 220658.s.at | Figure 3886: DNA330451, NP_110429.1, 221249_s_at |
| Figure 3836: PRO83844 | Figure 3887: PRO85652 |
| Figure 3837: DNA329307, NP_037483.1, 220684_at | Figure 3888: DNA330452, NP_112494.2, 221258_s_at Figure 3889: PRO85653 |
| Figure 3838: PRO84896 | Figure 3890: DNA295327, NP_068575.1, 221271_at |
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| Figure 3840: PRO80512 | Figure 3892: DNA330453, NP_112597.1, 221277_s_at |
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| Figure 3901: PRO85656 | Figure 3955: DNA325039, NP_004902.1, 221824_s_at |
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| Figure 3907: PRO84904 | Figure 3962: PRO85663 |
| Figure 3908: DNA330456, NP_060571.1, 221520_s_at | Figure 3963A-B: DNA330465, 253695.2, 221916_at |
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| Figure 3933: PRO49875 | Figure 3988: PRO69490 |
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| Figure 3935: PRO50083 | Figure 3990: PRO51716 |
| Figure 3936: DNA330460, NP_060255.2, 221685_s_at | Figure 3991: DNA328977, NP_071344.1, 222216_s_at |
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| Figure 3938: DNA273185, DNA273185, 221727_at | Figure 3993: DNA330469, NP_056249.1, 222250_s_at |
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| Figure 4007: DNA331607, HSA251830, 222392_x_at | Figure 4060: DNA327942, NP_060596.1, 222642_s_at |
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| Figure 4219: DNA327954, NP_113646.1, 223220_s_at | Figure 4272: PRO84939 |
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| Figure 4220: PRO83879 | Figure 4273: DNA330531, NP_037508.1, 223394_at |
| Figure 4221: DNA330515, NP_004580.1, 223221_at | Figure 4274: PRO85718 |
| Figure 4222: PRO85705 | Figure 4275: DNA329361, AF161528, 223397_s_at |
| Figure 4223: DNA329321, SEC13L, 223225_s_at | Figure 4276: PRO84940 |
| Figure 4224: PRO84906 | Figure 4277: DNA324156, NP_115588.1, 223403_s_at |
| Figure 4225: DNA247474, NP_054895.1, 223229_at | Figure 4278: PRO80856 |
| Figure 4226: PRO44999 | Figure 4279A-B: DNA254516, Clorf25, 223404_s_at |
| Figure 4227: DNA330516, AK000796, 223239_at | Figure 4280: PRO49623 |
| Figure 4228: PRO85706 | Figure 4281: DNA256407, NP_055188.1, 223423_at |
| Figure 4229: DNA287171, NP_036312.1, 223240_at | Figure 4282: PRO51448 |
| Figure 4230: PRO69462 | Figure 4283: DNA255676, HSM801648, 223434_at |
| Figure 4231: DNA324046, NP_115700.1, 223272_s_at | Figure 4284: PRO50738 |
| Figure 4232: PRO80763 | Figure 4285: DNA330532, NP_078804.1, 223439_at |
| Figure 4233: DNA330517, NP_115879.1, 223273_at Figure 4234: PRO85707 | Figure 4286: PRO85719 Figure 4287: DNA330533, NP_058647.1, 223451_s_at |
| Figure 4235: DNA330518, BC002493, 223274_at | Figure 4288: PRO772 |
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| Figure 4239: DNA330520, NP_005777.2, 223283_s_at | Figure 4292: PRO23554 |
| Figure 4240: PRO85710 | Figure 4293: DNA329456, NP_057126.1, 223489_x_at |
| Figure 4241: DNA330521, BC002762, 223286_at | Figure 4294: PRO85023 |
| Figure 4242: PRO85711 | Figure 4295: DNA329456, RRP40, 223490_s_at |
| Figure 4243A-B: DNA330522, AF250920, | Figure 4296: PRO85023 |
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| Figure 4244: PRO85712 | Figure 4298: PRO85720 |
| Figure 4245: DNA330523, BC001220, 223294_at | Figure 4299: DNA304784, NP_006564.1, 223502_s_at, |
| Figure 4246: PRO85713 | Figure 4300: PRO738 |
| Figure 4247: DNA330524, MGC4268, 223297_at | Figure 4301: DNA330535, NP_115883.1, 223506_at |
| Figure 4248: PRO85714 | Figure 4302: PRO85721 |
| Figure 4249: DNA329356, NP_115671.1, 223304_at | Figure 4303: DNA330536, NP_115666.1, 223542_at |
| Figure 4250: PRO84935 | Figure 4304: PRO85722 |
| Figure 4251: DNA330454, BC002551, 223307_at | Figure 4305: DNA330537, AF155827, 223556_at |
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| Figure 4253: DNA330526, NP_115682.1, 223318_s_at | Figure 4307A-B: DNA327908, HSM801808, |
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| Figure 4256: PRO85716 | • |
| Figure 4257: DNA329358, NP_115649.1, 223334_at Figure 4258: PRO84937 | Figure 4310: PRO85723 Figure 4311: DNA330539, NP_055411.1, 223639_s_at |
| Figure 4259: DNA330528, AF151063, 223335_at | Figure 4312: PRO85724 |
| Figure 4260: PRO50764 | Figure 4313: DNA330540, NP_055081.1, 223640_at |
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| Figure 4263A-B: DNA255756, HUMPDE7A, | Figure 4316: PRO85726 |
| 223358_s_at | Figure 4317: DNA330542, NP_115493.1, 223700_at |
| Figure 4264: PRO50812 | Figure 4318: PRO85727 |
| Figure 4265: DNA227125, AF132297, 223377_x_at | Figure 4319: DNA330543, NAG73, 223725_at |
| Figure 4266: PRO37588 | Figure 4320: PRO85728 |
| Figure 4267: DNA331614, CDCA1, 223381_at | Figure 4321: DNA329367, TTYH2, 223741_s_at |
| Figure 4268: PRO38881 | Figure 4322: PRO84946 |
| Figure 4269A-B: DNA329360, NP_115644.1, | Figure 4323: DNA331615, AB049635, 223743_s_at |
| 223382_s_at | Figure 4324: PRO62669 |
| Figure 4270: PRO84939 | Figure 4325: DNA188735, NP_001506.1, 223758_s_at |
| Figure 4271A-B: DNA329360, NIN283, 223383_at | Figure 4326: PRO26224 |
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| Figure 4327: DNA287253, LOC85028, 223774_at | Figure 4380: DNA330556, NP_061881.2, 224319_s_at |
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| Figure 4328: PRO69527 | Figure 4381: PRO85739 |
| Figure 4329: DNA331616, BC004277, 223785_at | Figure 4382: DNA330557, C20orf154, 224320.s_at |
| Figure 4330: PRO86613 | Figure 4383: PRO85740 |
| Figure 4331: DNA330544, NP_277049.1, 223800_s_at | Figure 4384: DNA330558, NP_057588.1, 224330_s_at |
| Figure 4332: PRO85729 | Figure 4385: PRO84950 |
| Figure 4333: DNA256005, NP_004842.1, 223806_s_at | Figure 4386: DNA327949, MRP64, 224334_s_at |
| Figure 4334: PRO51056 | Figure 4387: PRO83874 |
| Figure 4335: DNA330545, AF233516, 223834_at | Figure 4388A-B: DNA330559, BAB21791.1, |
| Figure 4336: PRO70906 | 224336_s_at |
| Figure 4337: DNA327200, NP_114156.1, 223836_at | Figure 4389: PRO85741 |
| Figure 4338: PRO1065 | Figure 4390: DNA331619, BC010896, 224345_x_at |
| Figure 4339: DNA330546, AF132203, 223839_s_at | Figure 4391: PRO86616 |
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| Figure 4341: DNA330547, NP_066014.1, 223849_s_at Figure 4342: PRO85731 | Figure 4393: PRO86617 |
| Figure 4343: DNA331392, NP_004186.1, 223851_s_at | Figure 4394: DNA272626, RIP5, 224376_s_at Figure 4395: PRO60759 |
| Figure 4344: PRO364 | Figure 4395: PNO30739 Figure 4396: DNA330560, NP_510882.1, 224413_s_at |
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| Figure 4347A-B: DNA330522, FOXP1, 223936_s_at | Figure 4399: PRO85743 |
| Figure 4348: PRO85712 | Figure 4400: DNA328323, NP_114148.2, 224428_s_at |
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| 223946_at | Figure 4402: DNA331621, AF060225, 224437_s_at |
| Figure 4350: PRO85734 | Figure 4403: PRO86618 |
| Figure 4351: DNA331393, D83532, 223961_s_at | Figure 4404: DNA330562, NP_115716.1, 224448_s_at |
| Figure 4352: PRO86458 | Figure 4405: PRO85744 |
| Figure 4353: DNA324248, SP110, 223980_s_at | Figure 4406: DNA330563, NP_113668.1, 224450_s_at |
| Figure 4354: PRO80932 | Figure 4407: PRO85745 |
| Figure 4355: DNA330551, BC009946, 223983_s_at | Figure 4408: DNA330564, NP_115885.1, 224451_x_at |
| Figure 4356: PRO85735 | Figure 4409: PRO85746 |
| Figure 4357: DNA330552, BC001104, 223984_s_at | Figure 4410: DNA330565, BC006111, 224454_at |
| Figure 4358: PRO85736 | Figure 4411: PRO85747 |
| Figure 4359: DNA328847, NP_056338.1, 223989_s_at | Figure 4412: DNA330566, NP_115720.1, 224464_s_at |
| Figure 4360: PRO84579 | Figure 4413: PRO85748 |
| Figure 4361: DNA331617, AF332652, 224046_s_at | Figure 4414: DNA329373, NP_115722.1, 224467_s_at |
| Figure 4362: PRO86614 | Figure 4415: PRO84952 |
| Figure 4363: DNA329369, AF293026, 224130_s_at Figure 4364: PRO84948 | Figure 4416: DNA330567, NP_116114.1, 224504_s_at |
| | Figure 4417: PRO85749 Figure 4418: DNA 227076 NP 116120 1 224511 c at |
| Figure 4365: DNA330553, AF116653, 224148.at Figure 4366: DNA331618, AF231339, 224204_x_at | Figure 4418: DNA327976, NP_116120.1, 224511_s_at Figure 4419: PRO69574 |
| Figure 4367: PRO86615 | Figure 4420: DNA330568, BC006428, 224516_s_at |
| Figure 4368: DNA330554, AF277993, 224211_at | Figure 4421: PRO85750 |
| Figure 4369: PRO85737 | Figure 4422: DNA329374, NP_115735.1, 224523_s_at |
| Figure 4370A-C: DNA227619, NP_054831.1, | Figure 4423: PRO84953 |
| 224218_s_at | Figure 4424: DNA331622, TNFRSF18, 224553.s.at |
| Figure 4371: PRO38082 | Figure 4425: PRO86619 |
| Figure 4372: DNA324707, NP_037369.1, 224232_s_at | Figure 4426: DNA330569, BC020516, 224572_s_at |
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| Figure 4374: DNA323935, NP_060586.1, 224233_s_at | Figure 4428: DNA330571, AK027320, 224607_s_at |
| Figure 4375: PRO80668 | Figure 4429: PRO85752 |
| Figure 4376: DNA329370, NP_060611.2, 224247_s_at | Figure 4430: DNA327980, BC008959, 224615_x_at |
| Figure 4377: PRO84949 | Figure 4431: PRO83900 |
| Figure 4378A-B: DNA330555, HSM801768, | Figure 4432: DNA329376, BAA91036.1, 224632_at |
| 224308_s_at | Figure 4433: PRO84954 |
| Figure 4379: PRO85738 | Figure 4434: DNA330572, CAB82324.1, 224648_at |
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Figure 5187A-B: DNA330802, 7694410.1, 229686_at Figure 5241: PRO85035 Figure 5188: PRO85966 Figure 5242: DNA331671, 277648.15, 230375_at Figure 5189: DNA331668, 403448.4, 229699_at Figure 5243: PRO86666 Figure 5190: PRO86663 Figure 5244: DNA331672, 332195.1, 230391_at Figure 5191A-C: DNA330804, NP_055847.1, Figure 5245: PRO86667 229704_at Figure 5246: DNA257756, DNA257756, 230405_at Figure 5192: PRO85968 Figure 5247: DNA330823, 010867.1, 230449_x_at Figure 5193: DNA331669, 1466538.1, 229718_at Figure 5248: PRO85987 Figure 5249A-B: DNA331673, 333480.5, 230489_at Figure 5194: PRO86664 Figure 5195A-B: DNA227985, CBX6, 229733_s_at Figure 5250: PRO86668 Figure 5196: PRO38448 Figure 5251A-B: DNA330825, 406864.4, 230526_at Figure 5197: DNA330806, 200298.1, 229809_at Figure 5252: PRO85989 Figure 5198: PRO85970 Figure 5253: DNA331674, 059446.1, 230529_at Figure 5199: DNA330807, 334422.1, 229814_at Figure 5254: PRO86669 Figure 5200: PRO85971 Figure 5255: DNA330827, NP_079521.1, 230536_at Figure 5201: DNA328092, NP_002598.2, 229830_at Figure 5256: PRO85991 Figure 5202: PRO84003 Figure 5257: DNA330828, 233615.1, 230566_at Figure 5203: DNA330808, 1397087.3, 229838_at Figure 5258: PRO85992 Figure 5204: PRO85972 Figure 5259: DNA330829, 007717.1, 230580_at Figure 5205: DNA328972, BC009950, 229872_s_at Figure 5260: PRO85993 Figure 5206A-C: DNA330810, AF330041, 229881.at Figure 5261: DNA257789, NP_116219.1, 230656_s_at Figure 5207: PRO85974 Figure 5262: PRO52338 Figure 5208: DNA287290, AK001793, 229980_s_at Figure 5263: DNA330830, 216899.1, 230703_at Figure 5209: PRO69560 Figure 5264: PRO85994 Figure 5210: DNA330811, 1382987.2, 230000_at Figure 5265A-B: DNA328499, SORL1, 230707_at Figure 5211: PRO85975 Figure 5266: PRO84321 Figure 5212A-B: DNA330812, 000264.19, 230021_at Figure 5267A-B: DNA328099, 335889.1, 230779_at Figure 5213: PRO85976 Figure 5268: PRO84009 Figure 5214: DNA330813, 246201.1, 230036_at Figure 5269: DNA194391, HSM800477, 230848_s_at Figure 5215: PRO85977 Figure 5270: DNA330831, 208876.1, 230913_at Figure 5216: DNA258657, DNA258657, 230060_at Figure 5271: PRO85995 Figure 5217: PRO52596 Figure 5272: DNA330832, 253831.5, 230930_at Figure 5218: DNA330814, 309641.1, 230097_at Figure 5273: PRO85996 Figure 5219: PRO85978 Figure 5274: DNA304827, AF293462, 230966_at Figure 5220: DNA329467, 029236.1, 230110_at Figure 5275: PRO1265 Figure 5221: PRO85033 Figure 5276: DNA330833, 984179.1, 230970_at Figure 5222: DNA330815, AK057940, 230165_at Figure 5277: PRO85997 Figure 5223: PRO85979 Figure 5278: DNA331675, BC017477, 231094_at Figure 5224: DNA329468, BC011589, 230170_at Figure 5279: PRO86670 Figure 5280: DNA331676, 980781.1, 231109_at Figure 5225: PRO88 Figure 5226: DNA330816, 980409.1, 230192_at Figure 5281: PRO86671 Figure 5227: PRO85980 Figure 5282: DNA329473, 370473.13, 231124_x_at Figure 5228A-B: DNA194784, DNA194784, Figure 5283: PRO85038 230218_at Figure 5284: DNA330835, 399441.1, 231166_at Figure 5229: PRO24061 Figure 5285: PRO85999 Figure 5230: DNA331670, 373719.30, 230257_s_at Figure 5286A-B: DNA330836, 242968.17, 231169_at Figure 5231: PRO86665 Figure 5287: PRO86000 Figure 5232A-C: DNA330817, AB020335, 230265_at Figure 5288: DNA330837, 429490.1, 231182_at Figure 5233: PRO85981 Figure 5289: PRO86001 Figure 5234: DNA330818, 212282.1, 230304_at Figure 5290: DNA150808, HUMGBP1, 231577_s_at Figure 5235: PRO85982 Figure 5291: PRO12478 Figure 5236: DNA330819, 982802.1, 230337_at Figure 5292: DNA155700, DNA155700, 231579_s_at Figure 5237: PRO85983 Figure 5293: DNA330838, NP_037460.2, 231715_s_at Figure 5238: DNA330820, 230585.2, 230345_at Figure 5294: PRO80743 Figure 5239: PRO85984 Figure 5295: DNA330839, NP_060908.1, 231769_at Figure 5240: DNA329470, NP_002756.1, 230352_at Figure 5296: PRO86002

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| Figure 5310: PRO86006 | Figure 5364: DNA331683, 422960.1, 232504_at Figure 5365: PRO86678 |
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| Figure 5316: DNA331677, 981573.1, 231890.at | Figure 5370: PRO86031 |
| Figure 5317: PRO86672 | Figure 5371: DNA330869, 406591.1, 232687_at |
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| Figure 5323: PRO51311 | Figure 5378: PRO69661 |
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| Figure 5327: DNA330851, 337679.3, 232081_at | 233208_x_at |
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| Figure 5330: PRO86015 | Figure 5384: DNA330874, NP_057528.1, 233461_x_at |
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| Figure 5332: PRO86016 | Figure 5386: DNA331423, AF176071, 233467 s.at |
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| Figure 5340: PRO86674 | 233632_s_at |
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| Figure 5343: PRO86020 | Figure 5397: DNA329481, ASB2, 233857_s_at |
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| Figure 5345: PRO86021 | Figure 5399: DNA326800, XRN2, 233878_s_at |
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| Figure 5347: PRO86022 | Figure 5401: DNA330547, MOV10, 233917_s_at |
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| Figure 5349: PRO86675 | Figure 5403: DNA330878, NP_079111.1, 233937_at |
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| Figure 5408: PRO84916 | Figure 5464: DNA331690, 200228.1, 235199_at |
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| Figure 5410: PRO86680 | Figure 5466: DNA330896, 250896.1, 235213_at |
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| Figure 5414: PRO86682 | Figure 5470A-C: DNA330897, 332999.23, 235252_at |
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| Figure 5416: PRO86043 | Figure 5472: DNA324093, BC019263, 235256_s_at |
| Figure 5417: DNA330881, CRACC, 234306_s_at | Figure 5473: PRO80802 |
| Figure 5418: PRO1138 | Figure 5474: DNA260946, NP_115741.1, 235266_at |
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| Figure 5420: PRO84901 | Figure 5476A-C: DNA329379, 010205.2, 235287_at |
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| Figure 5422: PRO20110 | Figure 5478: DNA330898, 227608.1, 235299_at |
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| Figure 5424: PRO69682 | Figure 5480A-B: DNA330899, 7690822.1, 235306_at |
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| Figure 5427: DNA304813, NP_277053.1, 234973_at | 235331_x_at |
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| Figure 5432: PRO86045 | Figure 5487: PRO86063 |
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| Figure 5436: PRO51974 Figure 5437: DNA330885, AK055618, 235022_at | Figure 5492: DNA331692, 979330.2, 235425_at |
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| Figure 5459: PRO86054 | Figure 5514: DNA331695, 350462.1, 235652_at |
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| Figure 5519: PRO86077 | Figure 5575: PRO85052 |
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| Figure 5541: PRO86088 | Figure 5597: PRO86109 |
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| Figure 5547: PRO85050 | Figure 5603: PRO86112 |
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| Figure 5554: PRO85051 | Figure 5609: DNA330955, 311471.1, 238303_at |
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| Figure 5560: PRO86093 | |
| Figure 5561: DNA330933, 407881.1, 236506_at | Figure 5616: PRO9743 Figure 5617: DNA330957, 003538.1, 238509_at |
| Figure 5562: PRO86094 | O Company of the Comp |
| Figure 5563: DNA330934, 406491.1, 236595_at | Figure 5618: PRO86117 Figure 5619: DNA330958, 370339.1, 238541_at |
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| Figure 5568: PRO86097 | Figure 5623: DNA260158, DNA260158, 238551_at |
| Figure 5569: DNA330937, 205124.1, 236645_at | Figure 5624: DNA329495, 1447201.1, 238581_at |
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| Figure 5649: PRO86128 | Figure 5705: PRO86152 |
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| Figure 5653: PRO86697 | Figure 5709: PRO84857 |
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| Figure 5663: PRO86133 | Figure 5719: PRO86156 |
| Figure 5664: DNA330975, 207267.3, 239278_at | Figure 5720: DNA330999, 1454193.1, 240830_at |
| Figure 5665: PRO86134 | Figure 5721: PRO86157 |
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| Figure 5667: PRO86135 | Figure 5723: PRO86158 |
| Figure 5668: DNA329501, 205323.1, 239331_at | Figure 5724: DNA331704, CARS, 240983_s_at |
| Figure 5669: PRO85062 | Figure 5725: PRO86699 |
| Figure 5670: DNA330977, 996962.2, 239364_at | Figure 5726: DNA329504, 197187.1, 241365_at |
| Figure 5671: PRO86136 | Figure 5727: PRO85065 |
| Figure 5672: DNA330978, 039840.1, 239376_at | Figure 5728: DNA331001, 1499887.1, 241370_at |
| Figure 5673: PRO86137 | Figure 5729: PRO86159 |
| Figure 5674: DNA330979, 407730.7, 239388_at | Figure 5730: DNA331002, 231340.1, 241435_at |
| Figure 5675: PRO86138 | Figure 5731: PRO86160 |
| Figure 5676: DNA330980, 1134871.2, 239401_at | Figure 5732: DNA331003, 391185.30, 241495_at |
| Figure 5677: PRO86139 | Figure 5733: PRO86161 |
| Figure 5678: DNA330981, 406409.1, 239404_at | Figure 5734: DNA331004, 314498.1, 241505.at |
| Figure 5679: PRO86140 | Figure 5735: PRO86162 |
| Figure 5680: DNA330982, 980479.1, 239413_at | Figure 5736: DNA331005, 197725.1, 241722_x_at |
| Figure 5681: PRO86141 | Figure 5737: PRO86163 |
| Figure 5682: DNA328200, 405394.1, 239442_at | Figure 5738: DNA329505, BC017102, 241734_at |

| Figure 5739: DNA331006, 193718.1, 241740_at | Figure 5795: PRO86701 |
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| Figure 5740: PRO86164 | Figure 5796: DNA331707, 330870.5, 242625_at |
| Figure 5741: DNA331007, 405858.1, 241756_at | Figure 5797: PRO86702 |
| Figure 5742: PRO86165 | Figure 5798: DNA331030, 407930.2, 242648_at |
| Figure 5743: DNA331705, 428179.1, 241775_at | Figure 5799: PRO86188 |
| Figure 5744: PRO86700 | Figure 5800: DNA331031, 405967.1, 242669_at |
| Figure 5745: DNA 195721, DNA 195721, 241819_at | Figure 5801: PRO86189 |
| Figure 5746: DNA331009, 222011.1, 241824_at | Figure 5802A-C: DNA331708, NP_006258.2, |
| Figure 5747: PRO86167 | 242712_x_at |
| Figure 5748: DNA331010, 218800.1, 241843.at | Figure 5803: PRO86703 |
| Figure 5749: PRO86168 | Figure 5804A-C: DNA331033, AF330045, 242722.at |
| Figure 5750: DNA331011, 979953.1, 241859_at | Figure 5805: PRO86191 |
| Figure 5751: PRO86169 | Figure 5806: DNA331034, 7689086.1, 242735_x_at |
| Figure 5752: DNA331012, 030070.1, 241869_at | Figure 5807: PRO86192 |
| Figure 5753: PRO86170 | Figure 5808: DNA331035, 210512.1, 242783_at |
| Figure 5754: DNA331013, 406509.1, 241924_at | Figure 5809: PRO86193 |
| Figure 5755: PRO86171 | Figure 5810: DNA331036, 360991.1, 242836_at |
| Figure 5756: DNA329506, NP_387510.1, 241937_s_at | Figure 5811: PRO86194 |
| Figure 5757: PRO85067 | Figure 5812: DNA328224, 028975.1, 242859_at |
| Figure 5758: DNA331014, 1447958.2, 241985_at | Figure 5813: PRO84127 |
| Figure 5759: PRO86172 | Figure 5814: DNA331037, 206873.1, 242890_at |
| Figure 5760: DNA331015, 109159.1, 242031.at | Figure 5815: PRO86195 |
| Figure 5761: PRO86173 | Figure 5816: DNA331709, 017276.1, 242903_at |
| Figure 5762: DNA331016, 229438.1, 242051_at | Figure 5817: PRO86704 |
| Figure 5763: PRO86174 | Figure 5818: DNA331710, 227540.15, 242960_at |
| Figure 5764: DNA328213, 419856.5, 242059_at | Figure 5819: PRO86705 |
| Figure 5765: PRO84116 | Figure 5820: DNA331711, 427600.1, 243006_at |
| Figure 5766: DNA331017, 409906.7, 242060_x_at | Figure 5821: PRO86706 |
| Figure 5767: PRO86175 | Figure 5822: DNA331041, 982079.2, 243030_at |
| Figure 5768: DNA331018, 355930.1, 242110_at | Figure 5823: PRO86199 |
| Figure 5769: PRO86176 | Figure 5824: DNA331042, 019764.1, 243037_at |
| Figure 5770: DNA331019, 234788.2, 242245_at | Figure 5825: PRO86200 |
| Figure 5771: PRO86177 | Figure 5826: DNA331043, 005042.1, 243134_at |
| Figure 5772: DNA331020, 403459.1, 242261.at | Figure 5827: PRO86201 |
| Figure 5773: PRO86178 | Figure 5828: DNA331044, 226264.10, 243154_at |
| Figure 5774: DNA331021, 017309.1, 242268_at | Figure 5829: PRO86202 |
| Figure 5775: PRO86179 | Figure 5830: DNA331045, 066434.1, 243222_at |
| Figure 5776: DNA331022, BC009627, 242304_at | Figure 5831: PRO86203 |
| Figure 5777: DNA331023, 119753.1, 242362_at | Figure 5832: DNA331712, 005752.1, 243271_at |
| Figure 5778: PRO86181 | Figure 5833: PRO86707 |
| Figure 5779: DNA331024, 028992.1, 242388_x_at | Figure 5834: DNA331047, BC020624, 243362_s_at |
| Figure 5780: PRO86182 | Figure 5835: DNA331048, 7688599.1, 243366.s_at |
| Figure 5781: DNA328220, 239839.1, 242405_at | Figure 5836: PRO86206 |
| Figure 5782: PRO84123 | Figure 5837: DNA331049, 402027.4, 243395_at |
| Figure 5783: DNA331025, 127891.1, 242457_at | Figure 5838: PRO86207 |
| Figure 5784: PRO86183 | Figure 5839: DNA331713, 982999.2, 243423_at |
| Figure 5785: DNA328221, 221374.1, 242471_at | Figure 5840: PRO86708 |
| Figure 5786: PRO84124 | Figure 5841: DNA331051, 306804.1, 243469_at |
| Figure 5787: DNA257874, DNA257874, 242517_at | Figure 5842: PRO86209 |
| Figure 5788: DNA331026, 014632.1, 242518_at | Figure 5843: DNA331714, 332965.1, 243496_at |
| Figure 5789: PRO86184 | Figure 5844: PRO86709 |
| Figure 5790: DNA331027, 053796.1, 242560_at | Figure 5845: DNA331053, 243689.1, 243509_at |
| Figure 5791: PRO86185 | Figure 5846: PRO86211 |
| Figure 5792: DNA331028, 7693434.1, 242606_at | Figure 5847: DNA331715, 7683458.1, 243514_at |
| Figure 5793: PRO86186 | Figure 5848: PRO86710 |
| Figure 5794: DNA331706, 351474.1, 242617_at | Figure 5849: DNA331055, 1512996.3, 243561_at |
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| Figure 5850: PRO86213 | Figure 5904: PRO86712 |
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| Figure 5851: DNA258957, DNA258957, 243631_at | Figure 5905: DNA331718, AK024409, DNA92232_at |
| Figure 5852: DNA331056, 218946.1, 243759_at | Figure 5906: PRO86713 |
| Figure 5853: PRO86214 | Figure 5907: DNA96866, DNA96866, DNA96866_at |
| Figure 5854: DNA194184, DNA194184, 243764_at | Figure 5908: PRO6015 |
| Figure 5855: PRO23576 | Figure 5909: DNA331073, BC011775, DNA101926_at |
| Figure 5856: DNA331057, 031316.1, 243888_at | Figure 5910: PRO86229 |
| Figure 5857: PRO86215 | Figure 5911: DNA108670, DNA108670, |
| Figure 5858: DNA331058, 400813.1, 243918_at | DNA108670_at |
| Figure 5859: PRO86216 | Figure 5912: PRO7171 |
| Figure 5860: DNA331059, 035870.32, 243934_at | Figure 5913: DNA304467, BC004535, DNA108688_at |
| Figure 5861: PRO86217 | Figure 5914: PRO71043 |
| Figure 5862: DNA210271, DNA210271, 243999_at | Figure 5915A-B: DNA108728, DNA108728, |
| Figure 5863: PRO33803 | DNA108728_at |
| Figure 5864A-B: DNA331060, 406931.1, 244008_at | Figure 5916: PRO9741 |
| Figure 5865: PRO86218 | Figure 5917: DNA329215, ICOS, DNA108917_at |
| Figure 5866: DNA331061, 198683.4, 244026_at | Figure 5918: PRO7424 |
| Figure 5867: PRO86219 | Figure 5919: DNA331719, BC002424, DNA143288_at |
| Figure 5868: DNA331062, BC021973, 244052_at | Figure 5920: PRO12705 |
| Figure 5869: PRO23771 | Figure 5921A-B: DNA150956, HUMORFKG1P, |
| Figure 5870: DNA331716, 212607.1, 244267_at | DNA150956_at |
| Figure 5871: PRO86711 | Figure 5922: DNA330417, APOL6, DNA164989_at |
| Figure 5872: DNA331064, 006039.1, 244313_at | Figure 5923: PRO21341 |
| Figure 5873: PRO86221 | Figure 5924: DNA329483, AF384857, DNA166819_at |
| Figure 5874: DNA108738, DNA108738, 244321_at | Figure 5925: PRO20110 |
| Figure 5875: PRO9822 | Figure 5926: DNA26842, DNA26842, P_Z64949_at |
| Figure 5876: DNA331065, 341348.1, 244382.at | Figure 5927: PRO180 Figure 5928: DNA304468, NP_077300.1, P_Z93700_at |
| Figure 5877: PRO86222 | Figure 5929: PRO71044 |
| Figure 5878: DNA331066, 207228.1, 244443.at | Figure 5930: DNA39423, DNA39423, P_X52252_at |
| Figure 5879: PRO86223 | Figure 5931: PRO271 |
| Figure 5880: DNA328239, 331922.4, 244450.at Figure 5881: PRO84142 | Figure 5932: DNA330262, GW112, P.Z64962_at |
| Figure 5882: DNA331067, 164869.1, 244599_at | Figure 5933: PRO85493 |
| Figure 5883: PRO86224 | Figure 5934: DNA331074, AF252257, P_A37030_at |
| Figure 5884: DNA331068, 337465.1, 244677_at | Figure 5935: DNA60764, DNA60764, P_A46906_at |
| Figure 5885: PRO86225 | Figure 5936: PRO1265 |
| Figure 5886: DNA329512, 336575.1, 244780_at | Figure 5937: DNA331720, AF289594, P_A37063_at |
| Figure 5887: PRO85073 | Figure 5938: PRO86714 |
| Figure 5888: DNA331069, 008651.1, 244798_at | Figure 5939: DNA331721, BC017876, P_A37079_at |
| Figure 5889: PRO86226 | Figure 5940: PRO71045 |
| Figure 5890: DNA331070, 393412.1, 244801_at | Figure 5941: DNA76401, DNA76401, P_A37126_at |
| Figure 5891: PRO86227 | Figure 5942: PRO1575 |
| Figure 5892: DNA331071, 343563.1, 244869.at | Figure 5943: DNA304475, NP_116246.1, P_A37128_at |
| Figure 5893: PRO86228 | Figure 5944: PRO71049 |
| Figure 5894A-B: DNA254566, BAA11502.1, | Figure 5945: DNA66480, HSAPO1, NM_000043_at |
| D80007_at | Figure 5946: PRO1207 |
| Figure 5895: PRO49669 | Figure 5947: DNA88195, CD3G, NM_000073_at |
| Figure 5896: DNA328961, BC011049, DNA36995_at | Figure 5948: PRO2693 |
| Figure 5897: PRO84667 | Figure 5949: DNA325712, CDK4, NM_000075_at |
| Figure 5898A-B: DNA331072, AB046821, | Figure 5950: PRO82194 |
| DNA53991_at | Figure 5951: DNA329934, BC013083, NM_000099_at |
| Figure 5899: DNA327200, KSP37, DNA59602_at | Figure 5952: PRO2721 |
| Figure 5900: PRO1065 | Figure 5953A-B: DNA331722, HUMFVA, |
| Figure 5901: DNA327205, GBP5, DNA61875_at | NM_000130_at |
| Figure 5902: PRO83478 | Figure 5954: PRO36374 |
| Figure 5903: DNA331717, BC020203, DNA71289_at | Figure 5955: DNA331723, U66095, NM_000161_at |
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Figure 5956: PRO86715 Figure 5999: PRO65 Figure 5957: DNA227668, HUMGLYKINB, Figure 6000A-B: DNA331732, HSCCR5AB2, NM_000167_at NM_000579_at Figure 5958: PRO38131 Figure 6001: PRO25194 Figure 5959A-D: DNA331724, HSGLA, Figure 6002: DNA290585, NP_000573.1, NM_000169_at NM_000582_f_at Figure 5960: DNA331725, BC006342, NM_000175_at Figure 6003: PRO70536 Figure 5961: DNA150823, NP_000185.1, Figure 6004: DNA216500, NP_000575.1, NM_000194_at NM_000584_at Figure 5962: PRO12810 Figure 6005: PRO34252 Figure 5963: DNA331726, HUMICAMA1A, Figure 6006: DNA36712, HUMIL3, NM_000588_at NM_000201_at Figure 6007: PRO67 Figure 5964: PRO86716 Figure 6008A-B: DNA331733, AF361105, Figure 5965A-B: DNA88419, HSINTA6R, NM_000590_at NM_000210_at Figure 6009: DNA331734, BC014081, NM_000593_at Figure 5966: PRO2339 Figure 6010: PRO36996 Figure 5967: DNA88428, HUMLAP, NM_000211_at Figure 6011A-B: DNA331735, AY066019, Figure 5968: PRO2787 NM_000594_at Figure 5969: DNA226014, NP_000230.1, Figure 6012A-B: DNA331736, AY070490, NM_000239_at NM_000595_at Figure 5970: PRO36477 Figure 6013: DNA331737, BC009902, NM_000597_at Figure 5971: DNA97287, NP_000240.1, Figure 6014: PRO2587 NM_000249_at Figure 6015: DNA217246, NP_000591.1, Figure 5972: PRO3634 NM_000600_at Figure 5973: DNA88554, NP_000241.1, Figure 6016: PRO34288 NM_000250_at Figure 6017: DNA331075, NP_000601.2, Figure 5974: PRO2839 NM_000610_at Figure 5975: DNA331727, BC008015, NM_000269_at Figure 6018: PRO86231 Figure 5976: PRO37534 Figure 6019A-C: DNA331738, AF375790, Figure 5977A-E: DNA331728, PTEN4, NM_000314_at NM_000619_at Figure 5978: DNA83020, NP_000349.1, Figure 6020A-B: DNA220752, ITGAM, NM_000358_at NM_000632_at Figure 5979: PRO2561 Figure 6021: PRO34730 Figure 5980: DNA227081, EGR2, NM_000399_at Figure 6022A-B: DNA97288, HUMBCL2C, Figure 5981: PRO37544 NM_000633_at Figure 5982: DNA76512, HSIL2REC, NM_000417_at Figure 6023: PRO3635 Figure 5983: PRO2020 Figure 6024: DNA331739, A12178, NM_000636_at Figure 5984: DNA76514, HSIL4R, NM_000418_at Figure 6025: PRO86720 Figure 5985: PRO2540 Figure 6026: DNA331740, HUMHPC, NM_000639_at Figure 5986: DNA329522, NP_000433.2, Figure 6027: PRO1208 NM_000442_at Figure 6028: DNA329000, HSU03905, NM_000647_at Figure 5987: PRO85080 Figure 6029: PRO84690 Figure 5988: DNA188732, NP_000475.1, Figure 6030: DNA328253, NP_004029.1, NM_000484_at NM_000699_at Figure 5989: PRO25302 Figure 6031: PRO84149 Figure 5990: DNA331729, AF281258, NM_000517_at Figure 6032: DNA89242, ANXA1, NM_000700_at Figure 5991: DNA331730, BC014514, NM_000527_at Figure 6033: PRO2907 Figure 5992: PRO2915 Figure 6034: DNA88194, CD3E, NM_000733_at Figure 5993: DNA331731, HSASM2MR, Figure 6035: PRO2220 NM_000543_at Figure 6036: DNA329975, PRO2325, NM_000791_at Figure 5994: DNA76516, IL6R, NM_000565_at Figure 6037: DNA331741, BC003097, NM_000873_at Figure 5995: PRO2022 Figure 6038: PRO86721 Figure 5996: DNA36718, HUMIL10, NM_000572_at Figure 6039: DNA331076, HSIFNABR, Figure 5997: PRO73 NM_000874_at Figure 5998: DNA324158, NP_000567.1, Figure 6040: PRO86232 NM_000576_at Figure 6041A-B: DNA83101, NP_000868.1,

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Figure 6132: PRO86234 Figure 6179: DNA227014, NP_002190.1, NM_002199_at Figure 6133: DNA331752, BC010240, NM_001908_at Figure 6180: PRO37477 Figure 6134: PRO86727 Figure 6181A-B: DNA88427, HSFNRB, Figure 6135: DNA225810, HSCATHL, NM_001912_at Figure 6136: PRO36273 NM_002211_at Figure 6182: PRO2786 Figure 6137: DNA83048, DEFA4, NM_001925_at Figure 6183: DNA103215, NP_002210.1, Figure 6138: PRO2057 Figure 6139: DNA88215, NP_001919.1, NM_002219_at NM_001928_at Figure 6184: PRO4545 Figure 6185: DNA331758, S82269, NM_002222_at Figure 6140: PRO2703 Figure 6186: PRO86731 Figure 6141: DNA196562, HSPCHDP7, Figure 6187: DNA331759, BC002646, NM_002228_at NM_001935_at Figure 6142: PRO25042 Figure 6188: PRO4671 Figure 6189: DNA331760, BC009466, NM_002229_at Figure 6143: DNA226871, NP_001942.1, Figure 6190: PRO4650 NM_001951_at Figure 6191A-B: DNA331761, AF305731S5, Figure 6144: PRO37334 NM_002250_at Figure 6145: DNA227332, NP_001943.1, Figure 6192: DNA150971, KLRB1, NM_002258_at NM_001952_at Figure 6193: PRO12564 Figure 6146: PRO37795 Figure 6194: DNA326343, BC003572, NM_002265_at Figure 6147: DNA225661, ECGF1, NM_001953_at Figure 6195: PRO82739 Figure 6148: PRO36124 Figure 6196: DNA288243, LAG3, NM_002286_at Figure 6149: DNA273174, HSEF1DELA, Figure 6197: PRO36451 NM_001960_at Figure 6198A-B: DNA188301, LIF, NM_002309_at Figure 6150: PRO61211 Figure 6199: PRO21834 Figure 6151: DNA150779, HUMETR103, Figure 6200A-B: DNA331762, HUMLYTOXBB, NM_001964_at NM_002341_at Figure 6152: PRO12798 Figure 6201: DNA88666, NP_002334.1, Figure 6153: DNA331753, HUMENOG, NML002343_at NM_001975_at Figure 6202: PRO2892 Figure 6154: PRO38010 Figure 6203: DNA227150, LY6E, NM_002346_at Figure 6155: DNA331754, BC002464, NM_001992_at Figure 6204: PRO37613 Figure 6156: PRO86728 Figure 6205: DNA327255, BC001061, NM_002394_at Figure 6157: DNA331755, D83920, NM_002003_at Figure 6206: PRO57298 Figure 6158: PRO86729 Figure 6207: DNA150937, HSU94352, NM_002405_at Figure 6159: DNA226881, HUMERGBFLI, Figure 6208: PRO11598 NM_002017_at Figure 6209: DNA82376, CXCL9, NM_002416_at Figure 6160: PRO37344 Figure 6210: PRO1723 Figure 6161: DNA88332, FVT1, NM_002035_at Figure 6211: DNA103283, MNDA, NM_002432_at Figure 6162: PRO2753 Figure 6212: PRO4613 Figure 6163: DNA225979, G1P3, NM_002038_at Figure 6213: DNA103525, NP_002457.1, Figure 6164: PRO36442 NM_002466_at Figure 6165: DNA331756, BC002666, NM_002053_at Figure 6214: PRO4852 Figure 6166: PRO12478 Figure 6215A-B: DNA331763, AF058696, Figure 6167: DNA88374, GZMK, NM_002104_at Figure 6168: PRO2768 NM_002485_at Figure 6169: DNA228014, ICAM3, NM_002162_at Figure 6216: PRO36001 Figure 6217: DNA103382, HSU49395, NM_002561_at Figure 6170: PRO38477 Figure 6171: DNA331757, A17548, NM_002167_at Figure 6218: PRO4711 Figure 6219A-B: DNA88331, HSFUR, NM_002569_at Figure 6172: PRO86730 Figure 6173: DNA76517, IL7R, NM_002185_at Figure 6220: PRO2752 Figure 6221: DNA103488, PCNA, NM_002592_at Figure 6174: PRO2541 Figure 6222: PRO4815 Figure 6175: DNA188271, NP_002179.1, Figure 6223: DNA328587, NP_002612.1, NM_002188_at NM_002621_at Figure 6176: PRO21795 Figure 6224: PRO2854 Figure 6177: DNA226396, IL15RA, NM_002189_at Figure 6225: DNA331764, NP_071438.1, Figure 6178: PRO36859

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Figure 6589: PRO86240 Figure 6630: DNA 196569, NP_056957.1, Figure 6590: DNA227233, NP_055157.1, NM_015873_at Figure 6631: PRO19859 NM_014342_at Figure 6632: DNA150865, LOC51596, NM_015921_at Figure 6591: PRO37696 Figure 6592: DNA227351, AF191020, NM_014367_at Figure 6633: PRO11587 Figure 6634: DNA150832, NP_057019.2, Figure 6593: PRO37814 NML015935_at Figure 6594: DNA331088, NP-055252.2, NM_014437_at Figure 6635: PRO12491 Figure 6595: PRO80674 Figure 6636: DNA331089, NP_057143.1, Figure 6596: DNA330084, SIT, NM_014450_at NM_016059_at Figure 6597: PRO9895 Figure 6637: PRO4984 Figure 6638: DNA331818, AF151899, NM_016072_at Figure 6598: DNA324198, NP_055400.1, Figure 6639: PRO793 NM_014585_at Figure 6640: DNA328663, AK001280, NM_016073_at Figure 6599: PRO37675 Figure 6641: PRO36183 Figure 6600A-B: DNA151879, NP_055463.1, Figure 6642: DNA331819, BC006807, NM_016077_at NM_014648_at Figure 6643: PRO38080 Figure 6601: PRO12743 Figure 6644: DNA150661, LOC51030, NM_016078_at Figure 6602: DNA194805, NP_055500.1, Figure 6645: PRO12398 NM_014685_at Figure 6646: DNA329292, AF085360, NM_016101_at Figure 6603: PRO24075 Figure 6647: PRO84882 Figure 6604A-B: DNA150467, AB018335, Figure 6648: DNA329923, HSPC035, NM_016127_at NM_014698_at Figure 6649: PRO85237 Figure 6605: PRO12272 Figure 6650: DNA304832, NP_057327.1, Figure 6606A-B: DNA194778, KIAA0152, NM_016243_at NM_014730_at Figure 6651: PRO71239 Figure 6607: PRO24056 Figure 6652: DNA328831, AF126780, NM_016245_at Figure 6608A-B: DNA277809, KIAA0275, Figure 6653: PRO233 NM_014767_at Figure 6654: DNA328513, AF151895, NM_016283_at Figure 6609: PRO64556 Figure 6655: PRO37815 Figure 6610A-B: DNA227353, SEC24D, Figure 6656: DNA304781, LOC51184, NM_016301_at NM_014822_at Figure 6657: PRO71191 Figure 6611: PRO37816 Figure 6658: DNA331820, BC001144, NM_016306_at Figure 6612: DNA93507, NP_055694.1, Figure 6659: PRO1080 NM_014879_at Figure 6660: DNA331821, AK023410, NM_016354_at Figure 6613: PRO4948 Figure 6661: PRO86752 Figure 6614A-B: DNA150954, KIAA0022, Figure 6662: DNA330390, AF178985, NM_016546_at NM_014880_at Figure 6663: PRO85599 Figure 6615: PRO12558 Figure 6664: DNA331822, AF318357, NM_016553_at Figure 6616A-B: DNA227293, DNA227293, Figure 6665: PRO86753 NM_014883_at Figure 6666: DNA227298, NP_057649.1, Figure 6617: PRO37756 NM_016565_at Figure 6618: DNA150805, FAM3C, NM_014888_at Figure 6667: PRO37761 Figure 6619: PRO11583 Figure 6668: DNA327869, NRN1, NM_016588_at Figure 6620A-B: DNA194837, NP_055714.1, Figure 6669: PRO1898 NM_014899_at Figure 6670: DNA331823, AK027682, NM_017424_at Figure 6621: PRO24100 Figure 6622A-B: DNA304464, CHSY1, Figure 6671: PRO86754 Figure 6672: DNA225694, FLJ20005, NM_017617_at NM_014918_at Figure 6673: PRO36157 Figure 6623: PRO71042 Figure 6674: DNA326385, NP_060117.2, Figure 6624: DNA330103, MD-2, NM_015364_at NM_017647_at Figure 6625: PRO19671 Figure 6626: DNA150872, NP_056202.1, Figure 6675: PRO82778 Figure 6676: DNA287206, FLJ20073, NM_017654_at NM_015387_at Figure 6677: PRO69488 Figure 6627: PRO12814 Figure 6628: DNA328590, BC001232, NM_015864_at Figure 6678: DNA227294, FLJ20303, NM_017755_at Figure 6679: PRO37757 Figure 6629: PRO84375

Figure 6680: DNA226646, NP_060352.1,

NM_017882_at

Figure 6681: PRO37109

Figure 6682: DNA331824, BC010907, NM_017906_at

Figure 6683: PRO86755

Figure 6684: DNA330537, HELLS, NM_018063_at

Figure 6685: PRO81892

Figure 6686: DNA328628, BC011983, NM_018072_at

Figure 6687: PRO84406

Figure 6688: DNA328841, BC003082, NM_018087_at

Figure 6689: PRO84575

Figure 6730: DNA327199, DJ971N18.2,

NM_021156_at

Figure 6731: PRO83475

Figure 6732: DNA227276, NP_005702.1,

NML021618_at

Figure 6733: PRO37739

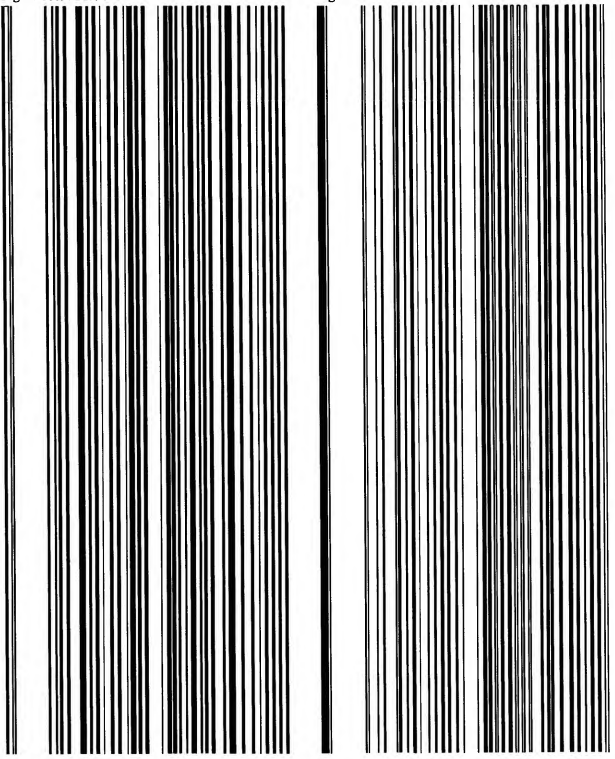
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Figure 6735: PRO86758

Figure 6736: DNA331833, AF269133, NM_021798_at

Figure 6737: PRO86759



HUMKG1BB_at Figure 6822: PRO82665 Figure 6823A-B: DNA287330, AB032991, Figure 6776: PRO86762 AB032991_at Figure 6777A-B: DNA331842, BC004375, Figure 6824: DNA331848, 1510819.1, P_X99863_at AF261758_at Figure 6825: PRO86764 Figure 6778: PRO38492 Figure 6826: DNA331849, HSINSP4BP, Figure 6779: DNA331095, NP_005216.1, HUME2F_at HSINSP4BP_at Figure 6780: PRO86245 Figure 6827: PRO66285 Figure 6781: DNA331843, AF202723, AB014568_at Figure 6782: DNA159542, DNA159542, Figure 6828A-B: DNA331850, HSA237724, HUMMAC30X_at HSA237724_at Figure 6829: DNA328049, 981676.1, HSM800856_at Figure 6783: DNA331844, BC009267, Figure 6830: PRO83963 **HUMLAMBBA_at** Figure 6831: DNA331851, AK027334, P_A51904_at Figure 6784: PRO82888 Figure 6785: DNA331096, S75881, P_V84330_at Figure 6832: PRO23392 Figure 6833: DNA193996, DNA193996, P_A40502_at Figure 6786: PRO86246 Figure 6834: PRO23400 Figure 6787: DNA287239, AF212242, AK024843 at Figure 6835: DNA194019, DNA194019, AK000004_at Figure 6788: PRO38497 Figure 6836: PRO23421 Figure 6789: DNA154390, DNA154390, Figure 6837: DNA194063, DNA194063, P_V84608_at HUMP13KIN_at Figure 6838: PRO23460 Figure 6790: DNA151247, DNA151247, P_V43601_at Figure 6839: DNA83046, NP_000565.1, P_X30170_at Figure 6791: PRO11643 Figure 6840: PRO2569 Figure 6792: DNA329950, MGC5576, P_V43613_at Figure 6841: DNA331852, 985629.1, P.Z59467.at Figure 6793: PRO11558 Figure 6842: PRO86765 Figure 6794: DNA161927, DNA161927, P.Z29229_at Figure 6843: DNA195915, DNA195915, P_X85020_at Figure 6795: DNA155316, DNA155316, P_A09058_at Figure 6796: DNA329026, AF230200, AK021966_at Figure 6844: DNA331853, BC001305, AK027031_at Figure 6845: PRO23769 Figure 6797A-B: DNA228052, DNA228052, Figure 6846A-B: DNA328720, NP_078800.2, AB006624_at P_X35729_at Figure 6798: PRO38515 Figure 6847: PRO84476 Figure 6799: DNA161913, DNA161913, Figure 6848: DNA194679, BAA05062.1, HSM800208_at HUMORFT_at Figure 6800: DNA331845, AK027432, Figure 6849: PRO23989 HSM800284_at Figure 6850: DNA331854, AF244129, AF244129_at Figure 6801: PRO86763 Figure 6851: PRO86766 Figure 6802: DNA329430, SPPL2A, AX027882_at Figure 6852: DNA 194766, DJ434O14.3, Figure 6803: PRO38524 HS434O143_at Figure 6804: DNA151422, DNA151422, P_X04312_at Figure 6853: PRO24046 Figure 6805: PRO11792 Figure 6854: DNA331855, BLP1, P.Z98236_at Figure 6806: DNA228066, NP_079431.1, Figure 6855: PRO85742 AK021910_at Figure 6856: DNA330358, BC008904, AX011617_at Figure 6807: PRO38529 Figure 6857: PRO85574 Figure 6808A-C: DNA330360, FYCO1, AK023397_at Figure 6858: DNA330380, FLJ12436, AK022498 at Figure 6809: PRO85576 Figure 6859: PRO85592 Figure 6810: DNA287185, DNA287185, P_V84564_at Figure 6811: PRO37492 Figure 6860: DNA328288, NP_073591.1, Figure 6812: DNA331846, AF272741, AK022938_at HUMTCBYY_at Figure 6861: PRO69876 Figure 6813: DNA331097, AK027322, AX041977_at Figure 6862: DNA196036, DNA196036, Figure 6814: PRO86247 AI471699_RC_at Figure 6863: DNA331098, AY052405, AX047348_at Figure 6815: DNA151756, DNA151756, P_X84947_at Figure 6864: PRO86248 Figure 6816: PRO12037 Figure 6865A-B: DNA331099, AB058685, Figure 6817: DNA151761, DNA151761, P_X84970_at AX048187_at Figure 6818: PRO12039 Figure 6866: DNA331100, BC021238, P_X84987_at Figure 6819: DNA331847, BC008330, AK026632_at Figure 6867: PRO86249 Figure 6820: PRO38556

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Figure 6868: DNA331101, NP_114143.1, 250446.2_at

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Figure 6915: PRO58324

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Figure 7317: PRO86796 Figure 7362: PRO63009 Figure 7318: DNA329047, BATF, NM_006399_at Figure 7363: DNA254157, HSU13045, NM_005254_at Figure 7364: PRO49271 Figure 7319: PRO58425 Figure 7365A-B: DNA124122, RBL2, NM_005611_at Figure 7320: DNA274167, AF026166, NM_006431_at Figure 7366: PRO6323 Figure 7321: PRO62097 Figure 7367: DNA330776, TOB1, NM_005749_at Figure 7322: DNA254572, NP_006576.1, Figure 7368: PRO58014 NM_006585_at Figure 7369: DNA326980, AF140598, NM_014248_at Figure 7323: PRO49675 Figure 7370: PRO83289 Figure 7324A-B: DNA331911, AB003334, Figure 7371: DNA271608, HUMRSC419, NM_006644_at NM_014670_at Figure 7325: PRO86797 Figure 7372: PRO59895 Figure 7326: DNA331912, BC009405, NM_013411_at Figure 7327: PRO86798 Figure 7373: DNA272928, HUMORFKG1F, Figure 7328: DNA255289, MELK, NM_014791_at NML014764_at Figure 7329: PRO50363 Figure 7374: PRO61012 Figure 7375: DNA290235, NP_057121.1, Figure 7330A-B: DNA331913, BAB21784.1, NM_016037_at NML015383_at Figure 7376: PRO70335 Figure 7331: PRO86799 Figure 7332: DNA329148, LOC51042, NM_015871_at Figure 7377: DNA331135, HUMKG1DD, HUMKG1DD_at Figure 7333: PRO84782 Figure 7378A-B: DNA330119, AF226044, Figure 7334: DNA326221, AF125098, NM_016095_at HUMKIAAQ_at Figure 7335: PRO82634 Figure 7379: PRO85381 Figure 7336: DNA331914, BC009398, Figure 7380: DNA331137, HS24P52, HUMP1CDC47_at HUMHSP70H_at Figure 7337: PRO86800 Figure 7338A-B: DNA328312, HUMAREB6, Figure 7381: PRO86278 Figure 7382A-B: DNA269805, NP_001263.1, HUMAREB6_at NM_001272_at Figure 7339: PRO84180 Figure 7383: PRO58209 Figure 7340: DNA325941, HSPCA, HSHSP90R_at Figure 7384: DNA270689, HSGATA3R, Figure 7341: PRO82388 NM_002051_at Figure 7342: DNA328483, VIT1, NM_000179_at Figure 7343: PRO84309 Figure 7385: PRO59053 Figure 7386: DNA331919, HUMCFA, NM_002965_at Figure 7344: DNA271847, HUMDNAJHOM, NM_001539_at Figure 7387: PRO80648 Figure 7345: PRO60127 Figure 7388A-B: DNA304800, NP_004146.1, NM_004155_at Figure 7346: DNA331915, BC001786, NM_002014_at Figure 7389: PRO69458 Figure 7347: PRO59262 Figure 7390: DNA273418, AAG01157.1, Figure 7348: DNA331916, HUMMIF, NM_002415_at Figure 7349: DNA331917, PHF1, NM_002636_at NM_004301_at Figure 7391: PRO61417 Figure 7350: PRO86802 Figure 7392: DNA330066, MLLT3, NM_004529_at Figure 7351: DNA329604, SRP54, NM_003136_at Figure 7393: PRO85346 Figure 7352: PRO85134 Figure 7394: DNA270733, S46622, NM_005605_at Figure 7353A-B: DNA331134, NP_003381.1, NM_003390_at Figure 7395: PRO59094 Figure 7354: PRO86275 Figure 7396: DNA331138, NP_005997.2, Figure 7355A-B: DNA290265, ZNF91, NM_006006_at NM_003430_f_at Figure 7397: PRO86279 Figure 7356: PRO70395 Figure 7398: DNA331139, NP_006865.1, Figure 7357A-C: DNA331918, AF009425, NM_006874_at NM_004338_at Figure 7399: PRO81172 Figure 7358: PRO86803 Figure 7400: DNA331920, AF090950, NM_015675_at Figure 7359: DNA254582, NP_004626.1, Figure 7401: PRO84384 Figure 7402: DNA329050, MRPS17, NM_015969_at NM_004635_at Figure 7403: PRO84712 Figure 7360: PRO49685 Figure 7361A-B: DNA275334, NP_112162.1, Figure 7404A-B: DNA329122, GS3955, NM_004749_at NM_021643_at

Figure 7452: PRO86283 Figure 7405: PRO84764 Figure 7453A-B: DNA331926, BAB13449.1, Figure 7406: DNA331921, 244055.1, AF320911_at AB046843_at Figure 7407: PRO86804 Figure 7454: PRO51258 Figure 7408: DNA331922, AK026275, AK026275_at Figure 7455: DNA255197, DNA255197, P.Z50392_at Figure 7409: PRO86805 Figure 7456: PRO50276 Figure 7410A-B: DNA254516, AF288399, Figure 7457: DNA328010, NP_149016.1, AF288399_at HSM801092_at Figure 7411: PRO49623 Figure 7458: PRO83928 Figure 7412: DNA328313, NP_115579.1, Figure 7459: DNA262805, DNA262805, AK025076_at HSM800425_at Figure 7413: PRO84181 Figure 7460: DNA331146, 1400830.1, Figure 7414: DNA327865, NP_079105.1, **HUMINLTRA_at** AK026315_at Figure 7461: PRO86284 Figure 7415: PRO83806 Figure 7462: DNA328317, cig5, AF026941_at Figure 7416: DNA294813, NP_444283.1, P_T67134_at Figure 7463: PRO69493 Figure 7417: PRO70763 Figure 7464: DNA331147, NP_079104.1, Figure 7418A-B: DNA254706, AB046851, AF131768_at AB046851_at Figure 7465: PRO86285 Figure 7419: DNA329052, NP_078801.1, Figure 7466: DNA255770, DNA255770, AK022106_at AK026237_at Figure 7467A-C: DNA254412, EVI5, AF008915_at Figure 7420: PRO84714 Figure 7468: PRO49522 Figure 7421: DNA256890, BC008988, P_Z00467_at Figure 7469: DNA331148, 978273.10, AK023244_at Figure 7422: PRO51824 Figure 7470: PRO86286 Figure 7423: DNA256291, FLJ21032, AK024685_f_at Figure 7471: DNA330532, AK026279, AK026279 at Figure 7424: PRO51335 Figure 7472: PRO85719 Figure 7425: DNA331923, HSUCP2X12, P_C69111_at Figure 7473: DNA330388, FLJ23468, AK027121_at Figure 7426: DNA213665, DNA213665, P.X30166_at Figure 7474: PRO85597 Figure 7427: PRO35126 Figure 7475: DNA331927, AK026969, AK026969_at Figure 7428: DNA331140, 332752.10, AK023798_at Figure 7476: PRO86807 Figure 7429: PRO86280 Figure 7477: DNA330447, FLJ22757, AK026410_at Figure 7430A-B: DNA331141, BAB13420.1, Figure 7478: PRO85648 AB046814_at Figure 7479: DNA324984, FLJ12298, AK022360 at Figure 7431: PRO86281 Figure 7480: PRO81578 Figure 7432: DNA331924, BC004932, AK024551_at Figure 7481: DNA331149, 7697327.1, HSM802839_at Figure 7433: PRO21434 Figure 7482: PRO86287 Figure 7434A-B: DNA256267, AB046838, Figure 7483A-B: DNA256267, DNA256267, AB046838_at AK023113_at Figure 7435: DNA327954, BAL, P_D00629_at Figure 7484: PRO51311 Figure 7436: PRO83879 Figure 7485: DNA331150, BC017725, 1387341.2_at Figure 7437: DNA255798, FLJ12377, AK022439_at Figure 7486: PRO86288 Figure 7438: PRO50853 Figure 7487: DNA257606, DNA257606, 428093.1 at Figure 7439: DNA330389, FLJ12888, AK022950_at Figure 7488: DNA258375, AF283301, 413231.5_at Figure 7440: PRO85598 Figure 7489: PRO52516 Figure 7441: DNA330086, FLJ12973, AK023035_at Figure 7490: DNA331928, AK027419, 154551.10 at Figure 7442: PRO85360 Figure 7491: PRO86808 Figure 7443: DNA331142, NP_116325.1, P_Z98137_at Figure 7492: DNA328319, BC019562, 411364.2 at Figure 7444: PRO51781 Figure 7493: DNA290812, DNA290812, Figure 7445: DNA329384, BC008502, P.Z33372_at 220495.3_CON_at Figure 7446: PRO84960 Figure 7494: PRO70559 Figure 7447A-B: DNA331143, NP_149075.2, Figure 7495: DNA304799, BC022410, 337588.1 at AK022613_at Figure 7496: PRO52633 Figure 7448: PRO86282 Figure 7497: DNA257403, DNA257403, 012814.1_at Figure 7449: DNA331925, 424693.10, AK022231_at Figure 7498: DNA304820, NP_115940.1, 317557.1_at Figure 7450: PRO86806 Figure 7499: PRO47351 Figure 7451: DNA331144, NP_078834.1, Figure 7500: DNA331929, BC019246, AK023982_at

441855.8_CON_at Figure 7545: DNA304817, BC015532, 211436.3 at Figure 7546: PRO71224 Figure 7501: PRO83338 Figure 7502: DNA260581, DNA260581, 127987.6_at Figure 7547: DNA260313, DNA260313, 1098929.1 at Figure 7503: PRO54507 Figure 7548: PRO54242 Figure 7504: DNA257576, DNA257576, 334945.2_at Figure 7549A-B: DNA328325, NP_061142.1, Figure 7505: DNA304819, BC004398, 202113.2_at 445188.4_at Figure 7506: DNA304794, FBXO30, 222128.1 at Figure 7550: PRO84190 Figure 7507: PRO71206 Figure 7551A-B: DNA304800, SERPINB9, Figure 7508: DNA259323, DNA259323, 022997.1_at 354740.1_at Figure 7509: PRO53256 Figure 7552: PRO69458 Figure 7510: DNA304796, MED8, 237428.13_at Figure 7553: DNA331156, 118180.1, 118180.1_at Figure 7511: PRO71208 Figure 7554: PRO86294 Figure 7512: DNA259615, DNA259615, 1000203.1_at Figure 7555: DNA287659, AK027790, 406833.1_at Figure 7513: DNA304805, AK027628, 475113.7_at Figure 7556: PRO69903 Figure 7514: PRO69531 Figure 7557: DNA331931, 092555.3, 092555.4_at Figure 7515: DNA304793, GBP4, 206425.2.at Figure 7558: PRO86810 Figure 7516: PRO71205 Figure 7559: DNA331157, NP_439896.1, 022541.5_at Figure 7560: PRO86295 Figure 7517: DNA331151, 018033.1, Figure 7561: DNA260573, DNA260573, 899597.1_at 018033.1_CON_at Figure 7562: PRO54499 Figure 7518: PRO86289 Figure 7563: DNA260157, DNA260157, 236833.1_at Figure 7519: DNA304068, AK057631, 1091656.1_at Figure 7564: PRO54086 Figure 7520: PRO71035 Figure 7521: DNA257714, EPSTI1, 337352.17_at Figure 7565: DNA174145, DNA174145, 100083.2.at Figure 7566: PRO35770 Figure 7522: PRO52268 Figure 7523: DNA304798, NP_443097.1, 246119.7_at Figure 7567: DNA260167, DNA260167, 264556.1_at Figure 7568A-B: DNA331932, 239260.1, 239260.1 at Figure 7524: PRO71210 Figure 7569: PRO86811 Figure 7525; DNA258721, DNA258721, 197627.1_at Figure 7526A-B: DNA257461, NP_113607.1, Figure 7570: DNA260031, DNA260031, 161526.1_at Figure 7571: DNA258907, DNA258907, 347940.2 at 086533.1_at Figure 7572: PRO52840 Figure 7527: PRO52040 Figure 7528: DNA331152, 1042156.3, 1042156.3 at Figure 7573: DNA257455, DNA257455, 977723.3 at Figure 7529: PRO86290 Figure 7574: PRO52035 Figure 7575: DNA304807, BC014978, 005415.2_at Figure 7530: DNA331153, 004052.1, 004052.1.at Figure 7576: PRO71216 Figure 7531: PRO86291 Figure 7577: DNA258864, DNA258864, 331965.1 at Figure 7532: DNA331930, AK054582, 978231.1_at Figure 7578: DNA304811, 428051.2, 428051.2.at Figure 7533: PRO86809 Figure 7534: DNA259587, DNA259587, 220866.1 at Figure 7579: PRO71220 Figure 7535: DNA106195, DNA106195, 359193.13.at Figure 7580: DNA257389, FLJ14906, 987098.1_at Figure 7536: DNA331154, 212376.1, 212376.1 at Figure 7581: PRO51974 Figure 7582: DNA331158, 130352.1, 130352.1_at Figure 7537: PRO86292 Figure 7538: DNA331155, 112652.1, 112652.1.at Figure 7583: PRO86296 Figure 7584: DNA258951, DNA258951, 222361.1_at Figure 7539: PRO86293 Figure 7540: DNA304806, BC019022, 983343.1_at Figure 7585: DNA331159, NP_077291.1, Figure 7541: PRO71215 411426.29_at Figure 7542: DNA262708, DNA262708, Figure 7586: PRO86297 118516.1_RC_at Figure 7587: DNA257784, DNA257784, 481853.1 at Figure 7543: DNA259475, DNA259475, 239162.1 at Figure 7588: DNA331933, AF272148, 074299.1 at Figure 7544: DNA269148, DNA269148, 411192.2_at Figure 7589: PRO86812

BRIEF DESCRIPTION OF THE DRAWINGS

In the list of figures for the present application, specific cDNA sequences which are differentially expressed in differentiated macrophages as compared to normal undifferentiated monocytes are individually identified with a specific alphanumerical designation. These cDNA sequences are differentially expressed in monocytes that are specifically treated as described in Example 1 below. If start and/or stop codons have been identified in a cDNA sequence shown in the attached figures, they are shown in bold and underlined font, and the encoded polypeptide is shown in the next consecutive figure.

The Figures 1-7589 show the nucleic acids of the invention and their encoded PRO polypeptides. Also included, for convenience is a List of Figures, which gives the figure number and the corresponding DNA or PRO number.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

I. <u>Definitions</u>

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The terms "PRO polypeptide" and "PRO" as used herein and when immediately followed by a numerical designation refer to various polypeptides, wherein the complete designation (i.e., PRO/number) refers to specific polypeptide sequences as described herein. The terms "PRO/number polypeptide" and "PRO/number" wherein the term "number" is provided as an actual numerical designation as used herein encompass native sequence polypeptides and polypeptide variants (which are further defined herein). The PRO polypeptides described herein may be isolated from a variety of sources, such as from human tissue types or from another source, or prepared by recombinant or synthetic methods. The term "PRO polypeptide" refers to each individual PRO/number polypeptide disclosed herein. All disclosures in this specification which refer to the "PRO polypeptide" refer to each of the polypeptides individually as well as jointly. For example, descriptions of the preparation of, purification of, derivation of, formation of antibodies to or against, administration of, compositions containing, treatment of a disease with, etc., pertain to each polypeptide of the invention individually. The term "PRO polypeptide" also includes variants of the PRO/number polypeptides disclosed herein.

A "native sequence PRO polypeptide" comprises a polypeptide having the same amino acid sequence as the corresponding PRO polypeptide derived from nature. Such native sequence PRO polypeptides can be isolated from nature or can be produced by recombinant or synthetic means. The term "native sequence PRO polypeptide" specifically encompasses naturally-occurring truncated or secreted forms of the specific PRO polypeptide (e.g., an extracellular domain sequence), naturally-occurring variant forms (e.g., alternatively spliced forms) and naturally-occurring allelic variants of the polypeptide. In various embodiments of the invention, the native sequence PRO polypeptides disclosed herein are mature or full-length native sequence polypeptides comprising the full-length amino acids sequences shown in the accompanying figures. Start and stop codons are shown in bold font and underlined in the figures. However, while the PRO polypeptide disclosed in the accompanying figures are shown to begin with methionine residues designated herein as amino acid position 1 in the figures, it is conceivable and possible that other methionine residues located either upstream or downstream from the amino acid position 1 in the figures may be employed as the starting amino acid residue for the PRO polypeptides.

The PRO polypeptide "extracellular domain" or "ECD" refers to a form of the PRO polypeptide which is essentially free of the transmembrane and cytoplasmic domains. Ordinarily, a PRO polypeptide

ECD will have less than 1% of such transmembrane and/or cytoplasmic domains and preferably, will have less than 0.5% of such domains. It will be understood that any transmembrane domains identified for the PRO polypeptides of the present invention are identified pursuant to criteria routinely employed in the art for identifying that type of hydrophobic domain. The exact boundaries of a transmembrane domain may vary but most likely by no more than about 5 amino acids at either end of the domain as initially identified herein. Optionally, therefore, an extracellular domain of a PRO polypeptide may contain from about 5 or fewer amino acids on either side of the transmembrane domain/extracellular domain boundary as identified in the Examples or specification and such polypeptides, with or without the associated signal peptide, and nucleic acid encoding them, are contemplated by the present invention.

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The approximate location of the "signal peptides" of the various PRO polypeptides disclosed herein are shown in the present specification and/or the accompanying figures. It is noted, however, that the C-terminal boundary of a signal peptide may vary, but most likely by no more than about 5 amino acids on either side of the signal peptide C-terminal boundary as initially identified herein, wherein the C-terminal boundary of the signal peptide may be identified pursuant to criteria routinely employed in the art for identifying that type of amino acid sequence element (e.g., Nielsen et al., Prot. Eng. 10:1-6 (1997) and von Heinje et al., Nucl. Acids. Res. 14:4683-4690 (1986)). Moreover, it is also recognized that, in some cases, cleavage of a signal sequence from a secreted polypeptide is not entirely uniform, resulting in more than one secreted species. These mature polypeptides, where the signal peptide is cleaved within no more than about 5 amino acids on either side of the C-terminal boundary of the signal peptide as identified herein, and the polynucleotides encoding them, are contemplated by the present invention.

"PRO polypeptide variant" means an active PRO polypeptide as defined above or below having at least about 80% amino acid sequence identity with a full-length native sequence PRO polypeptide sequence as disclosed herein, a PRO polypeptide sequence lacking the signal peptide as disclosed herein, an extracellular domain of a PRO polypeptide, with or without the signal peptide, as disclosed herein or any other fragment of a full-length PRO polypeptide sequence as disclosed herein. Such PRO polypeptide variants include, for instance, PRO polypeptides wherein one or more amino acid residues are added, or deleted, at the N- or C-terminus of the full-length native amino acid sequence. Ordinarily, a PRO polypeptide variant will have at least about 80% amino acid sequence identity, alternatively at least about 81% amino acid sequence identity, alternatively at least about 82% amino acid sequence identity, alternatively at least about 83% amino acid sequence identity, alternatively at least about 84% amino acid sequence identity, alternatively at least about 85% amino acid sequence identity, alternatively at least about 86% amino acid sequence identity, alternatively at least about 87% amino acid sequence identity, alternatively at least about 88% amino acid sequence identity, alternatively at least about 89% amino acid sequence identity, alternatively at least about 90% amino acid sequence identity, alternatively at least about 91% amino acid sequence identity, alternatively at least about 92% amino acid sequence identity, alternatively at least about 93% amino acid sequence identity, alternatively at least about 94% amino acid sequence identity, alternatively at least about 95% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 97% amino acid sequence identity, alternatively at least about 98% amino acid sequence identity and alternatively at least about 99% amino acid sequence identity to a full-length native sequence PRO polypeptide sequence as disclosed herein, a

PRO polypeptide sequence lacking the signal peptide as disclosed herein, an extracellular domain of a PRO polypeptide, with or without the signal peptide, as disclosed herein or any other specifically defined fragment of a full-length PRO polypeptide sequence as disclosed herein. Ordinarily, PRO variant polypeptides are at least about 10 amino acids in length, alternatively at least about 20 amino acids in length, alternatively at least about 40 amino acids in length, alternatively at least about 50 amino acids in length, alternatively at least about 60 amino acids in length, alternatively at least about 70 amino acids in length, alternatively at least about 80 amino acids in length, alternatively at least about 90 amino acids in length, alternatively at least about 100 amino acids in length, alternatively at least about 150 amino acids in length, alternatively at least about 200 amino acids in length, alternatively at least about 200 amino acids in length, alternatively at least about 300 amino acids in length, or more.

"Percent (%) amino acid sequence identity" with respect to the PRO polypeptide sequences identified herein is defined as the percentage of amino acid residues in a candidate sequence that are identical with the amino acid residues in the specific PRO polypeptide sequence, after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity, and not considering any conservative substitutions as part of the sequence identity. Alignment for purposes of determining percent amino acid sequence identity can be achieved in various ways that are within the skill in the art, for instance, using publicly available computer software such as BLAST, BLAST-2, ALIGN or Megalign (DNASTAR) software. Those skilled in the art can determine appropriate parameters for measuring alignment, including any algorithms needed to achieve maximal alignment over the full length of the sequences being compared. For purposes herein, however, % amino acid sequence identity values are generated using the sequence comparison computer program ALIGN-2, wherein the complete source code for the ALIGN-2 program is provided in Table 1 below. The ALIGN-2 sequence comparison computer program was authored by Genentech, Inc. and the source code shown in Table 1 below has been filed with user documentation in the U.S. Copyright Office, Washington D.C., 20559, where it is registered under U.S. Copyright Registration No. TXU510087. The ALIGN-2 program is publicly available through Genentech, Inc., South San Francisco, California or may be compiled from the source code provided in Table 1 below. The ALIGN-2 program should be compiled for use on a UNIX operating system, preferably digital UNIX V4.0D. All sequence comparison parameters are set by the ALIGN-2 program and do not vary.

In situations where ALIGN-2 is employed for amino acid sequence comparisons, the % amino acid sequence identity of a given amino acid sequence A to, with, or against a given amino acid sequence B (which can alternatively be phrased as a given amino acid sequence A that has or comprises a certain % amino acid sequence identity to, with, or against a given amino acid sequence B) is calculated as follows:

100 times the fraction X/Y

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where X is the number of amino acid residues scored as identical matches by the sequence alignment program ALIGN-2 in that program's alignment of A and B, and where Y is the total number of amino acid residues in B. It will be appreciated that where the length of amino acid sequence A is not equal to the length of amino acid sequence B, the % amino acid sequence identity of A to B will not equal the % amino acid sequence identity of B to A. As examples of % amino acid sequence identity calculations using this

method, Tables 2 and 3 demonstrate how to calculate the % amino acid sequence identity of the amino acid sequence designated "Comparison Protein" to the amino acid sequence designated "PRO", wherein "PRO" represents the amino acid sequence of a hypothetical PRO polypeptide of interest, "Comparison Protein" represents the amino acid sequence of a polypeptide against which the "PRO" polypeptide of interest is being compared, and "X, "Y" and "Z" each represent different hypothetical amino acid residues.

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Unless specifically stated otherwise, all % amino acid sequence identity values used herein are obtained as described in the immediately preceding paragraph using the ALIGN-2 computer program. However, % amino acid sequence identity values may also be obtained as described below by using the WU-BLAST-2 computer program (Altschul et al., Methods in Enzymology 266:460-480 (1996)). Most of the WU-BLAST-2 search parameters are set to the default values. Those not set to default values, i.e., the adjustable parameters, are set with the following values: overlap span = 1, overlap fraction = 0.125, word threshold (T) = 11, and scoring matrix = BLOSUM62. When WU-BLAST-2 is employed, a % amino acid sequence identity value is determined by dividing (a) the number of matching identical amino acid residues between the amino acid sequence of the PRO polypeptide of interest having a sequence derived from the native PRO polypeptide and the comparison amino acid sequence of interest (i.e., the sequence against which the PRO polypeptide of interest is being compared which may be a PRO variant polypeptide) as determined by WU-BLAST-2 by (b) the total number of amino acid residues of the PRO polypeptide of interest. For example, in the statement "a polypeptide comprising an the amino acid sequence A which has or having at least 80% amino acid sequence identity to the amino acid sequence B", the amino acid sequence A is the comparison amino acid sequence of interest and the amino acid sequence B is the amino acid sequence of the PRO polypeptide of interest.

Percent amino acid sequence identity may also be determined using the sequence comparison program NCBI-BLAST2 (Altschul et al., Nucleic Acids Res. 25:3389-3402 (1997)). The NCBI-BLAST2 sequence comparison program may be downloaded from http://www.ncbi.nlm.nih.gov or otherwise obtained from the National Institute of Health, Bethesda, MD. NCBI-BLAST2 uses several search parameters, wherein all of those search parameters are set to default values including, for example, unmask = yes, strand = all, expected occurrences = 10, minimum low complexity length = 15/5, multi-pass e-value = 0.01, constant for multi-pass = 25, dropoff for final gapped alignment = 25 and scoring matrix = BLOSUM62.

In situations where NCBI-BLAST2 is employed for amino acid sequence comparisons, the % amino acid sequence identity of a given amino acid sequence A to, with, or against a given amino acid sequence B (which can alternatively be phrased as a given amino acid sequence A that has or comprises a certain % amino acid sequence identity to, with, or against a given amino acid sequence B) is calculated as follows:

100 times the fraction X/Y

where X is the number of amino acid residues scored as identical matches by the sequence alignment program NCBI-BLAST2 in that program's alignment of A and B, and where Y is the total number of amino acid residues in B. It will be appreciated that where the length of amino acid sequence A is not equal to the

length of amino acid sequence B, the % amino acid sequence identity of A to B will not equal the % amino acid sequence identity of B to A.

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"PRO variant polynucleotide" or "PRO variant nucleic acid sequence" means a nucleic acid molecule which encodes an active PRO polypeptide as defined below and which has at least about 80% nucleic acid sequence identity with a nucleotide acid sequence encoding a full-length native sequence PRO polypeptide sequence as disclosed herein, a full-length native sequence PRO polypeptide sequence lacking the signal peptide as disclosed herein, an extracellular domain of a PRO polypeptide, with or without the signal peptide, as disclosed herein or any other fragment of a full-length PRO polypeptide sequence as disclosed herein. Ordinarily, a PRO variant polynucleotide will have at least about 80% nucleic acid sequence identity, alternatively at least about 81% nucleic acid sequence identity, alternatively at least about 82% nucleic acid sequence identity, alternatively at least about 83% nucleic acid sequence identity, alternatively at least about 84% nucleic acid sequence identity, alternatively at least about 85% nucleic acid sequence identity, alternatively at least about 86% nucleic acid sequence identity, alternatively at least about 87% nucleic acid sequence identity, alternatively at least about 88% nucleic acid sequence identity, alternatively at least about 89% nucleic acid sequence identity, alternatively at least about 90% nucleic acid sequence identity, alternatively at least about 91% nucleic acid sequence identity, alternatively at least about 92% nucleic acid sequence identity, alternatively at least about 93% nucleic acid sequence identity, alternatively at least about 94% nucleic acid sequence identity, alternatively at least about 95% nucleic acid sequence identity, alternatively at least about 96% nucleic acid sequence identity, alternatively at least about 97% nucleic acid sequence identity, alternatively at least about 98% nucleic acid sequence identity and alternatively at least about 99% nucleic acid sequence identity with a nucleic acid sequence encoding a fulllength native sequence PRO polypeptide sequence as disclosed herein, a full-length native sequence PRO polypeptide sequence lacking the signal peptide as disclosed herein, an extracellular domain of a PRO polypeptide, with or without the signal sequence, as disclosed herein or any other fragment of a full-length PRO polypeptide sequence as disclosed herein. Variants do not encompass the native nucleotide sequence.

Ordinarily, PRO variant polynucleotides are at least about 30 nucleotides in length, alternatively at least about 60 nucleotides in length, alternatively at least about 90 nucleotides in length, alternatively at least about 120 nucleotides in length, alternatively at least about 180 nucleotides in length, alternatively at least about 210 nucleotides in length, alternatively at least about 240 nucleotides in length, alternatively at least about 270 nucleotides in length, alternatively at least about 300 nucleotides in length, alternatively at least about 450 nucleotides in length, alternatively at least about 600 nucleotides in length, alternatively at least about 600 nucleotides in length, alternatively at least about 900 nucleotides in length, or more.

"Percent (%) nucleic acid sequence identity" with respect to PRO-encoding nucleic acid sequences identified herein is defined as the percentage of nucleotides in a candidate sequence that are identical with the nucleotides in the PRO nucleic acid sequence of interest, after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity. Alignment for purposes of determining percent nucleic acid sequence identity can be achieved in various ways that are within the skill in the art, for instance, using publicly available computer software such as BLAST, BLAST-2, ALIGN or Megalign (DNASTAR) software. For purposes herein, however, % nucleic acid sequence identity values are generated using the sequence comparison computer program ALIGN-2, wherein the complete source code

for the ALIGN-2 program is provided in Table 1 below. The ALIGN-2 sequence comparison computer program was authored by Genentech, Inc. and the source code shown in Table 1 below has been filed with user documentation in the U.S. Copyright Office, Washington D.C., 20559, where it is registered under U.S. Copyright Registration No. TXU510087. The ALIGN-2 program is publicly available through Genentech, Inc., South San Francisco, California or may be compiled from the source code provided in Table 1 below. The ALIGN-2 program should be compiled for use on a UNIX operating system, preferably digital UNIX V4.0D. All sequence comparison parameters are set by the ALIGN-2 program and do not vary.

In situations where ALIGN-2 is employed for nucleic acid sequence comparisons, the % nucleic acid sequence identity of a given nucleic acid sequence C to, with, or against a given nucleic acid sequence D (which can alternatively be phrased as a given nucleic acid sequence C that has or comprises a certain % nucleic acid sequence identity to, with, or against a given nucleic acid sequence D) is calculated as follows:

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100 times the fraction W/Z

where W is the number of nucleotides scored as identical matches by the sequence alignment program ALIGN-2 in that program's alignment of C and D, and where Z is the total number of nucleotides in D. It will be appreciated that where the length of nucleic acid sequence C is not equal to the length of nucleic acid sequence D, the % nucleic acid sequence identity of C to D will not equal the % nucleic acid sequence identity of D to C. As examples of % nucleic acid sequence identity calculations, Tables 4 and 5, demonstrate how to calculate the % nucleic acid sequence identity of the nucleic acid sequence designated "Comparison DNA" to the nucleic acid sequence designated "PRO-DNA", wherein "PRO-DNA" represents a hypothetical PRO-encoding nucleic acid sequence of interest, "Comparison DNA" represents the nucleotide sequence of a nucleic acid molecule against which the "PRO-DNA" nucleic acid molecule of interest is being compared, and "N", "L" and "V" each represent different hypothetical nucleotides.

Unless specifically stated otherwise, all % nucleic acid sequence identity values used herein are obtained as described in the immediately preceding paragraph using the ALIGN-2 computer program. However, % nucleic acid sequence identity values may also be obtained as described below by using the WU-BLAST-2 computer program (Altschul et al., Methods in Enzymology 266:460-480 (1996)). Most of the WU-BLAST-2 search parameters are set to the default values. Those not set to default values, i.e., the adjustable parameters, are set with the following values: overlap span = 1, overlap fraction = 0.125, word threshold (T) = 11, and scoring matrix = BLOSUM62. When WU-BLAST-2 is employed, a % nucleic acid sequence identity value is determined by dividing (a) the number of matching identical nucleotides between the nucleic acid sequence of the PRO polypeptide-encoding nucleic acid molecule of interest having a sequence derived from the native sequence PRO polypeptide-encoding nucleic acid and the comparison nucleic acid molecule of interest (i.e., the sequence against which the PRO polypeptide-encoding nucleic acid molecule of interest is being compared which may be a variant PRO polynucleotide) as determined by WU-BLAST-2 by (b) the total number of nucleotides of the PRO polypeptide-encoding nucleic acid molecule of interest. For example, in the statement "an isolated nucleic acid molecule comprising a nucleic acid sequence A which has or having at least 80% nucleic acid sequence identity to the nucleic acid sequence B", the nucleic acid sequence A is the comparison nucleic acid molecule of interest and the nucleic

acid sequence B is the nucleic acid sequence of the PRO polypeptide-encoding nucleic acid molecule of interest.

Percent nucleic acid sequence identity may also be determined using the sequence comparison program NCBI-BLAST2 (Altschul et al., Nucleic Acids Res. 25:3389-3402 (1997)). The NCBI-BLAST2 sequence comparison program may be downloaded from http://www.ncbi.nlm.nih.gov or otherwise obtained from the National Institute of Health, Bethesda, MD. NCBI-BLAST2 uses several search parameters, wherein all of those search parameters are set to default values including, for example, unmask = yes, strand = all, expected occurrences = 10, minimum low complexity length = 15/5, multi-pass e-value = 0.01, constant for multi-pass = 25, dropoff for final gapped alignment = 25 and scoring matrix = BLOSUM62.

In situations where NCBI-BLAST2 is employed for sequence comparisons, the % nucleic acid sequence identity of a given nucleic acid sequence C to, with, or against a given nucleic acid sequence D (which can alternatively be phrased as a given nucleic acid sequence C that has or comprises a certain % nucleic acid sequence identity to, with, or against a given nucleic acid sequence D) is calculated as follows:

100 times the fraction W/Z

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where W is the number of nucleotides scored as identical matches by the sequence alignment program NCBI-BLAST2 in that program's alignment of C and D, and where Z is the total number of nucleotides in D. It will be appreciated that where the length of nucleic acid sequence C is not equal to the length of nucleic acid sequence D, the % nucleic acid sequence identity of C to D will not equal the % nucleic acid sequence identity of D to C.

In other embodiments, PRO variant polynucleotides are nucleic acid molecules that encode an active PRO polypeptide and which are capable of hybridizing, preferably under stringent hybridization and wash conditions, to nucleotide sequences encoding a full-length PRO polypeptide as disclosed herein. PRO variant polypeptides may be those that are encoded by a PRO variant polynucleotide.

"Isolated," when used to describe the various polypeptides disclosed herein, means polypeptide that has been identified and separated and/or recovered from a component of its natural environment. Contaminant components of its natural environment are materials that would typically interfere with diagnostic or therapeutic uses for the polypeptide, and may include enzymes, hormones, and other proteinaceous or non-proteinaceous solutes. In preferred embodiments, the polypeptide will be purified (1) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence by use of a spinning cup sequenator, or (2) to homogeneity by SDS-PAGE under non-reducing or reducing conditions using Coomassie blue or, preferably, silver stain. Isolated polypeptide includes polypeptide in situ within recombinant cells, since at least one component of the PRO polypeptide natural environment will not be present. Ordinarily, however, isolated polypeptide will be prepared by at least one purification step.

An "isolated" PRO polypeptide-encoding nucleic acid or other polypeptide-encoding nucleic acid is a nucleic acid molecule that is identified and separated from at least one contaminant nucleic acid molecule with which it is ordinarily associated in the natural source of the polypeptide-encoding nucleic acid. An isolated polypeptide-encoding nucleic acid molecule is other than in the form or setting in which it is found in nature. Isolated polypeptide-encoding nucleic acid molecules therefore are distinguished from the

specific polypeptide-encoding nucleic acid molecule as it exists in natural cells. However, an isolated polypeptide-encoding nucleic acid molecule includes polypeptide-encoding nucleic acid molecules contained in cells that ordinarily express the polypeptide where, for example, the nucleic acid molecule is in a chromosomal location different from that of natural cells.

The term "control sequences" refers to DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

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Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, "operably linked" means that the DNA sequences being linked are contiguous, and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist, the synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice.

The term "antibody" is used in the broadest sense and specifically covers, for example, single anti-PRO monoclonal antibodies (including agonist, antagonist, and neutralizing antibodies), anti-PRO antibody compositions with polyepitopic specificity, single chain anti-PRO antibodies, and fragments of anti-PRO antibodies (see below). The term "monoclonal antibody" as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies, i.e., the individual antibodies comprising the population are identical except for possible naturally-occurring mutations that may be present in minor amounts.

"Stringency" of hybridization reactions is readily determinable by one of ordinary skill in the art, and generally is an empirical calculation dependent upon probe length, washing temperature, and salt concentration. In general, longer probes require higher temperatures for proper annealing, while shorter probes need lower temperatures. Hybridization generally depends on the ability of denatured DNA to reanneal when complementary strands are present in an environment below their melting temperature. The higher the degree of desired homology between the probe and hybridizable sequence, the higher the relative temperature which can be used. As a result, it follows that higher relative temperatures would tend to make the reaction conditions more stringent, while lower temperatures less so. For additional details and explanation of stringency of hybridization reactions, see Ausubel et al., Current Protocols in Molecular Biology, Wiley Interscience Publishers, (1995).

"Stringent conditions" or "high stringency conditions", as defined herein, may be identified by those that: (1) employ low ionic strength and high temperature for washing, for example 0.015 M sodium chloride/0.0015 M sodium citrate/0.1% sodium dodecyl sulfate at 50°C; (2) employ during hybridization a denaturing agent, such as formamide, for example, 50% (v/v) formamide with 0.1% bovine serum albumin/0.1% Ficoll/0.1% polyvinylpyrrolidone/50mM sodium phosphate buffer at pH 6.5 with 750 mM sodium chloride, 75 mM sodium citrate at 42°C; or (3) employ 50% formamide, 5 x SSC (0.75 M NaCl,

0.075 M sodium citrate), 50 mM sodium phosphate (pH 6.8), 0.1% sodium pyrophosphate, 5 x Denhardt's solution, sonicated salmon sperm DNA (50 µg/ml), 0.1% SDS, and 10% dextran sulfate at 42°C, with washes at 42°C in 0.2 x SSC (sodium chloride/sodium citrate) and 50% formamide at 55°C, followed by a high-stringency wash consisting of 0.1 x SSC containing EDTA at 55°C.

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"Moderately stringent conditions" may be identified as described by Sambrook et al., Molecular Cloning: A Laboratory Manual, New York: Cold Spring Harbor Press, 1989, and include the use of washing solution and hybridization conditions (e.g., temperature, ionic strength and %SDS) less stringent that those described above. An example of moderately stringent conditions is overnight incubation at 37°C in a solution comprising: 20% formamide, 5 x SSC (150 mM NaCl, 15 mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5 x Denhardt's solution, 10% dextran sulfate, and 20 mg/ml denatured sheared salmon sperm DNA, followed by washing the filters in 1 x SSC at about 37-50°C. The skilled artisan will recognize how to adjust the temperature, ionic strength, etc. as necessary to accommodate factors such as probe length and the like.

The term "epitope tagged" when used herein refers to a chimeric polypeptide comprising a PRO polypeptide fused to a "tag polypeptide". The tag polypeptide has enough residues to provide an epitope against which an antibody can be made, yet is short enough such that it does not interfere with activity of the polypeptide to which it is fused. The tag polypeptide preferably also is fairly unique so that the antibody does not substantially cross-react with other epitopes. Suitable tag polypeptides generally have at least six amino acid residues and usually between about 8 and 50 amino acid residues (preferably, between about 10 and 20 amino acid residues).

As used herein, the term "immunoadhesin" designates antibody-like molecules which combine the binding specificity of a heterologous protein (an "adhesin") with the effector functions of immunoglobulin constant domains. Structurally, the immunoadhesins comprise a fusion of an amino acid sequence with the desired binding specificity which is other than the antigen recognition and binding site of an antibody (i.e., is "heterologous"), and an immunoglobulin constant domain sequence. The adhesin part of an immunoadhesin molecule typically is a contiguous amino acid sequence comprising at least the binding site of a receptor or a ligand. The immunoglobulin constant domain sequence in the immunoadhesin may be obtained from any immunoglobulin, such as IgG-1, IgG-2, IgG-3, or IgG-4 subtypes, IgA (including IgA-1 and IgA-2), IgE, IgD or IgM.

"Active" or "activity" for the purposes herein refers to form(s) of a PRO polypeptide which retain a biological and/or an immunological activity of native or naturally-occurring PRO, wherein "biological" activity refers to a biological function (either inhibitory or stimulatory) caused by a native or naturally-occurring PRO other than the ability to induce the production of an antibody against an antigenic epitope possessed by a native or naturally-occurring PRO and an "immunological" activity refers to the ability to induce the production of an antibody against an antigenic epitope possessed by a native or naturally-occurring PRO.

The term "antagonist" is used in the broadest sense, and includes any molecule that partially or fully blocks, inhibits, or neutralizes a biological activity of a native PRO polypeptide disclosed herein. In a similar manner, the term "agonist" is used in the broadest sense and includes any molecule that mimics a biological activity of a native PRO polypeptide disclosed herein. Suitable agonist or antagonist molecules

specifically include agonist or antagonist antibodies or antibody fragments, fragments or amino acid sequence variants of native PRO polypeptides, peptides, antisense oligonucleotides, small organic molecules, etc. Methods for identifying agonists or antagonists of a PRO polypeptide may comprise contacting a PRO polypeptide with a candidate agonist or antagonist molecule and measuring a detectable change in one or more biological activities normally associated with the PRO polypeptide.

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"Treatment" refers to both therapeutic treatment and prophylactic or preventative measures, wherein the object is to prevent or slow down (lessen) the targeted pathologic condition or disorder. Those in need of treatment include those already with the disorder as well as those prone to have the disorder or those in whom the disorder is to be prevented.

"Chronic" administration refers to administration of the agent(s) in a continuous mode as opposed to an acute mode, so as to maintain the initial therapeutic effect (activity) for an extended period of time. "Intermittent" administration is treatment that is not consecutively done without interruption, but rather is cyclic in nature.

"Mammal" for purposes of treatment refers to any animal classified as a mammal, including humans, domestic and farm animals, and zoo, sports, or pet animals, such as dogs, cats, cattle, horses, sheep, pigs, goats, rabbits, etc. Preferably, the mammal is human.

Administration "in combination with" one or more further therapeutic agents includes simultaneous (concurrent) and consecutive administration in any order.

"Carriers" as used herein include pharmaceutically acceptable carriers, excipients, or stabilizers which are nontoxic to the cell or mammal being exposed thereto at the dosages and concentrations employed. Often the physiologically acceptable carrier is an aqueous pH buffered solution. Examples of physiologically acceptable carriers include buffers such as phosphate, citrate, and other organic acids; antioxidants including ascorbic acid; low molecular weight (less than about 10 residues) polypeptide; proteins, such as serum albumin, gelatin, or immunoglobulins; hydrophilic polymers such as polyvinylpyrrolidone; amino acids such as glycine, glutamine, asparagine, arginine or lysine; monosaccharides, disaccharides, and other carbohydrates including glucose, mannose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sorbitol; salt-forming counterions such as sodium; and/or nonionic surfactants such as TWEENTM, polyethylene glycol (PEG), and PLURONICSTM.

"Antibody fragments" comprise a portion of an intact antibody, preferably the antigen binding or variable region of the intact antibody. Examples of antibody fragments include Fab, Fab', F(ab')₂, and Fv fragments; diabodies; linear antibodies (Zapata et al., <u>Protein Eng.</u> 8(10): 1057-1062 [1995]); single-chain antibody molecules; and multispecific antibodies formed from antibody fragments.

Papain digestion of antibodies produces two identical antigen-binding fragments, called "Fab" fragments, each with a single antigen-binding site, and a residual "Fc" fragment, a designation reflecting the ability to crystallize readily. Pepsin treatment yields an F(ab')₂ fragment that has two antigen-combining sites and is still capable of cross-linking antigen.

"Fv" is the minimum antibody fragment which contains a complete antigen-recognition and - binding site. This region consists of a dimer of one heavy- and one light-chain variable domain in tight, non-covalent association. It is in this configuration that the three CDRs of each variable domain interact to define an antigen-binding site on the surface of the V_{H} - V_{L} dimer. Collectively, the six CDRs confer antigen-

binding specificity to the antibody. However, even a single variable domain (or half of an Fv comprising only three CDRs specific for an antigen) has the ability to recognize and bind antigen, although at a lower affinity than the entire binding site.

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The Fab fragment also contains the constant domain of the light chain and the first constant domain (CH1) of the heavy chain. Fab fragments differ from Fab' fragments by the addition of a few residues at the carboxy terminus of the heavy chain CH1 domain including one or more cysteines from the antibody hinge region. Fab'-SH is the designation herein for Fab' in which the cysteine residue(s) of the constant domains bear a free thiol group. F(ab')₂ antibody fragments originally were produced as pairs of Fab' fragments which have hinge cysteines between them. Other chemical couplings of antibody fragments are also known.

The "light chains" of antibodies (immunoglobulins) from any vertebrate species can be assigned to one of two clearly distinct types, called kappa and lambda, based on the amino acid sequences of their constant domains.

Depending on the amino acid sequence of the constant domain of their heavy chains, immunoglobulins can be assigned to different classes. There are five major classes of immunoglobulins: IgA, IgD, IgE, IgG, and IgM, and several of these may be further divided into subclasses (isotypes), e.g., IgG1, IgG2, IgG3, IgG4, IgA, and IgA2.

"Single-chain Fv" or "sFv" antibody fragments comprise the V_H and V_L domains of antibody, wherein these domains are present in a single polypeptide chain. Preferably, the Fv polypeptide further comprises a polypeptide linker between the V_H and V_L domains which enables the sFv to form the desired structure for antigen binding. For a review of sFv, see Pluckthun in <u>The Pharmacology of Monoclonal Antibodies</u>, vol. 113, Rosenburg and Moore eds., Springer-Verlag, New York, pp. 269-315 (1994).

The term "diabodies" refers to small antibody fragments with two antigen-binding sites, which fragments comprise a heavy-chain variable domain (V_H) connected to a light-chain variable domain (V_L) in the same polypeptide chain $(V_{H^-}V_L)$. By using a linker that is too short to allow pairing between the two domains on the same chain, the domains are forced to pair with the complementary domains of another chain and create two antigen-binding sites. Diabodies are described more fully in, for example, EP 404,097; WO 93/11161; and Hollinger et al., <u>Proc. Natl. Acad. Sci. USA</u>, 90:6444-6448 (1993).

An "isolated" antibody is one which has been identified and separated and/or recovered from a component of its natural environment. Contaminant components of its natural environment are materials which would interfere with diagnostic or therapeutic uses for the antibody, and may include enzymes, hormones, and other proteinaceous or nonproteinaceous solutes. In preferred embodiments, the antibody will be purified (1) to greater than 95% by weight of antibody as determined by the Lowry method, and most preferably more than 99% by weight, (2) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence by use of a spinning cup sequenator, or (3) to homogeneity by SDS-PAGE under reducing or nonreducing conditions using Coomassie blue or, preferably, silver stain. Isolated antibody includes the antibody in situ within recombinant cells since at least one component of the antibody's natural environment will not be present. Ordinarily, however, isolated antibody will be prepared by at least one purification step.

An antibody that "specifically binds to" or is "specific for" a particular polypeptide or an epitope on a particular polypeptide is one that binds to that particular polypeptide or epitope on a particular polypeptide without substantially binding to any other polypeptide or polypeptide epitope.

The word "label" when used herein refers to a detectable compound or composition which is conjugated directly or indirectly to the antibody so as to generate a "labeled" antibody. The label may be detectable by itself (e.g. radioisotope labels or fluorescent labels) or, in the case of an enzymatic label, may catalyze chemical alteration of a substrate compound or composition which is detectable.

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By "solid phase" is meant a non-aqueous matrix to which the antibody of the present invention can adhere. Examples of solid phases encompassed herein include those formed partially or entirely of glass (e.g., controlled pore glass), polysaccharides (e.g., agarose), polyacrylamides, polystyrene, polyvinyl alcohol and silicones. In certain embodiments, depending on the context, the solid phase can comprise the well of an assay plate; in others it is a purification column (e.g., an affinity chromatography column). This term also includes a discontinuous solid phase of discrete particles, such as those described in U.S. Patent No. 4,275,149.

A "liposome" is a small vesicle composed of various types of lipids, phospholipids and/or surfactant which is useful for delivery of a drug (such as a PRO polypeptide or antibody thereto) to a mammal. The components of the liposome are commonly arranged in a bilayer formation, similar to the lipid arrangement of biological membranes.

A "small molecule" is defined herein to have a molecular weight below about 500 Daltons.

The term "immune related disease" means a disease in which a component of the immune system of a mammal causes, mediates or otherwise contributes to a morbidity in the mammal. Also included are diseases in which stimulation or intervention of the immune response has an ameliorative effect on progression of the disease. Included within this term are immune-mediated inflammatory diseases, non-immune-mediated inflammatory diseases, infectious diseases, immunodeficiency diseases, neoplasia, etc.

The term "T cell mediated disease" means a disease in which T cells directly or indirectly mediate or otherwise contribute to a morbidity in a mammal. The T cell mediated disease may be associated with cell mediated effects, lymphokine mediated effects, etc., and even effects associated with B cells if the B cells are stimulated, for example, by the lymphokines secreted by T cells.

Examples of immune-related and inflammatory diseases, some of which are immune or T cell mediated, which can be treated according to the invention include systemic lupus erythematosis, rheumatoid arthritis, juvenile chronic arthritis, spondyloarthropathies, systemic sclerosis (scleroderma), idiopathic inflammatory myopathies (dermatomyositis, polymyositis), Sjögren's syndrome, systemic vasculitis, sarcoidosis, autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria), autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia), thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis), diabetes mellitus, immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis), demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy, hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis,

granulomatous hepatitis, and sclerosing cholangitis, inflammatory bowel disease (ulcerative colitis: Crohn's disease), gluten-sensitive enteropathy, and Whipple's disease, autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis, allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria, immunologic diseases of the lung such as eosinophilic pneumonias, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis, transplantation associated diseases including graft rejection and graft -versus-host-disease. Infectious diseases including viral diseases such as AIDS (HIV infection), hepatitis A, B, C, D, and E, herpes, etc., bacterial infections, fungal infections, protozoal infections and parasitic infections.

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The term "effective amount" is a concentration or amount of a PRO polypeptide and/or agonist/antagonist which results in achieving a particular stated purpose. An "effective amount" of a PRO polypeptide or agonist or antagonist thereof may be determined empirically. Furthermore, a "therapeutically effective amount" is a concentration or amount of a PRO polypeptide and/or agonist/antagonist which is effective for achieving a stated therapeutic effect. This amount may also be determined empirically.

The term "cytotoxic agent" as used herein refers to a substance that inhibits or prevents the function of cells and/or causes destruction of cells. The term is intended to include radioactive isotopes (e.g., I¹³¹, I¹²⁵, Y⁹⁰ and Re¹⁸⁶), chemotherapeutic agents, and toxins such as enzymatically active toxins of bacterial, fungal, plant or animal origin, or fragments thereof.

A "chemotherapeutic agent" is a chemical compound useful in the treatment of cancer. Examples of chemotherapeutic agents include adriamycin, doxorubicin, epirubicin, 5-fluorouracil, cytosine arabinoside ("Ara-C"), cyclophosphamide, thiotepa, busulfan, cytoxin, taxoids, e.g., paclitaxel (Taxol, Bristol-Myers Squibb Oncology, Princeton, NJ), and doxetaxel (Taxotere, Rhône-Poulenc Rorer, Antony, France), toxotere, methotrexate, cisplatin, melphalan, vinblastine, bleomycin, etoposide, ifosfamide, mitomycin C, mitoxantrone, vincristine, vinorelbine, carboplatin, teniposide, daunomycin, carminomycin, aminopterin, dactinomycin, mitomycins, esperamicins (see U.S. Pat. No. 4,675,187), melphalan and other related nitrogen mustards. Also included in this definition are hormonal agents that act to regulate or inhibit hormone action on tumors such as tamoxifen and onapristone.

A "growth inhibitory agent" when used herein refers to a compound or composition which inhibits growth of a cell, especially cancer cell overexpressing any of the genes identified herein, either in vitro or in vivo. Thus, the growth inhibitory agent is one which significantly reduces the percentage of cells overexpressing such genes in S phase. Examples of growth inhibitory agents include agents that block cell cycle progression (at a place other than S phase), such as agents that induce G1 arrest and M-phase arrest. Classical M-phase blockers include the vincas (vincristine and vinblastine), taxol, and topo II inhibitors such as doxorubicin, epirubicin, daunorubicin, etoposide, and bleomycin. Those agents that arrest G1 also spill over into S-phase arrest, for example, DNA alkylating agents such as tamoxifen, prednisone, dacarbazine, mechlorethamine, cisplatin, methotrexate, 5-fluorouracil, and ara-C. Further information can be found in The Molecular Basis of Cancer, Mendelsohn and Israel, eds., Chapter 1, entitled "Cell cycle regulation, oncogens, and antineoplastic drugs" by Murakami et al. (WB Saunders: Philadelphia, 1995), especially p. 13.

The term "cytokine" is a generic term for proteins released by one cell population which act on another cell as intercellular mediators. Examples of such cytokines are lymphokines, monokines, and

traditional polypeptide hormones. Included among the cytokines are growth hormone such as human growth hormone, N-methionyl human growth hormone, and bovine growth hormone; parathyroid hormone; thyroxine; insulin; proinsulin; relaxin; prorelaxin; glycoprotein hormones such as follicle stimulating hormone (FSH), thyroid stimulating hormone (TSH), and luteinizing hormone (LH); hepatic growth factor; fibroblast growth factor; prolactin; placental lactogen; tumor necrosis factor-α and -β; mullerian-inhibiting substance; mouse gonadotropin-associated peptide; inhibin; activin; vascular endothelial growth factor; integrin; thrombopoietin (TPO); nerve growth factors such as NGF-β; platelet-growth factor; transforming growth factors (TGFs) such as TGF-α and TGF-β; insulin-like growth factor-I and -II; erythropoietin (EPO); osteoinductive factors; interferons such as interferon-α, -β, and -γ, colony stimulating factors (CSFs) such as macrophage-CSF (M-CSF); granulocyte-macrophage-CSF (GM-CSF); and granulocyte-CSF (G-CSF); interleukins (ILs) such as IL-1, IL-1α, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-11, IL-12; a tumor necrosis factor such as TNF-α or TNF-β; and other polypeptide factors including LIF and kit ligand (KL). As used herein, the term cytokine includes proteins from natural sources or from recombinant cell culture and biologically active equivalents of the native sequence cytokines.

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As used herein, the term "immunoadhesin" designates antibody-like molecules which combine the binding specificity of a heterologous protein (an "adhesin") with the effector functions of immunoglobulin constant domains. Structurally, the immunoadhesins comprise a fusion of an amino acid sequence with the desired binding specificity which is other than the antigen recognition and binding site of an antibody (*i.e.*, is "heterologous"), and an immunoglobulin constant domain sequence. The adhesin part of an immunoadhesin molecule typically is a contiguous amino acid sequence comprising at least the binding site of a receptor or a ligand. The immunoglobulin constant domain sequence in the immunoadhesin may be obtained from any immunoglobulin, such as IgG-1, IgG-2, IgG-3, or IgG-4 subtypes, IgA (including IgA-1 and IgA-2), IgE, IgD or IgM.

As used herein, the term "inflammatory cells" designates cells that enhance the inflammatory response such as mononuclear cells, eosinophils, macrophages, and polymorphonuclear neutrophils (PMN).

Table 1

```
/*
5
       * C-C increased from 12 to 15
       * Z is average of EQ
       * B is average of ND
       * match with stop is _M; stop-stop = 0; J (joker) match = 0
10
                                    /* value of a match with a stop */
       #define _M
                  day[26][26] = {
       int
              ABCDEFGHIJKLMNOPQRSTUVWXYZ*/
                  { 2, 0,-2, 0, 0,-4, 1,-1,-1, 0,-1,-2,-1, 0,_M, 1, 0,-2, 1, 1, 0, 0,-6, 0,-3, 0},
       /* A */
15
                   { 0, 3,-4, 3, 2,-5, 0, 1,-2, 0, 0,-3,-2, 2,_M,-1, 1, 0, 0, 0, 0, 0,-2,-5, 0,-3, 1},
       /* B */
                    \{-2, -4, 15, -5, -5, -4, -3, -3, -2, 0, -5, -6, -5, -4, \underline{M}, -3, -5, -4, 0, -2, 0, -2, -8, 0, 0, -5\}, 
       /* C */
                   { 0, 3,-5, 4, 3,-6, 1, 1,-2, 0, 0,-4,-3, 2,_M,-1, 2,-1, 0, 0, 0,-2,-7, 0,-4, 2},
       /* D */
                   { 0, 2,-5, 3, 4,-5, 0, 1,-2, 0, 0,-3,-2, 1,_M,-1, 2,-1, 0, 0, 0,-2,-7, 0,-4, 3},
       /* E */
                   {-4,-5,-4,-6,-5, 9,-5,-2, 1, 0,-5, 2, 0,-4, M,-5,-5,-4,-3,-3, 0,-1, 0, 0, 7,-5},
20
       /* F */
                   \{1, 0, -3, 1, 0, -5, 5, -2, -3, 0, -2, -4, -3, 0, M, -1, -1, -3, 1, 0, 0, -1, -7, 0, -5, 0\},\
       /* G */
                   {-1, 1,-3, 1, 1,-2,-2, 6,-2, 0, 0,-2,-2, 2,_M, 0, 3, 2,-1,-1, 0,-2,-3, 0, 0, 2},
       /* H */
                   \{-1, -2, -2, -2, 1, -3, -2, 5, 0, -2, 2, 2, 2, -2, M, -2, -2, -1, 0, 0, 4, -5, 0, -1, -2\},\
       /* I */
                   /* J */
                   {-1, 0,-5, 0, 0,-5,-2, 0,-2, 0, 5,-3, 0, 1,_M,-1, 1, 3, 0, 0, 0,-2,-3, 0,-4, 0}
       /* K */
25
                   {-2,-3,-6,-4,-3, 2,-4,-2, 2, 0,-3, 6, 4,-3, M,-3,-2,-3,-3,-1, 0, 2,-2, 0,-1,-2}, {-1,-2,-5,-3,-2, 0,-3,-2, 2, 0, 0, 4, 6,-2, M,-2,-1, 0,-2,-1, 0, 2,-4, 0,-2,-1},
        /* L */
        /* M */
                   { 0, 2,-4, 2, 1,-4, 0, 2,-2, 0, 1,-3,-2, 2,_M,-1, 1, 0, 1, 0, 0,-2,-4, 0,-2, 1},
        /* N */
                   /* O */
                   \{1,-1,-3,-1,-1,-5,-1,0,-2,0,-1,-3,-2,-1,M,6,0,0,1,0,0,-1,-6,0,-5,0\},
        /* P */
30
                   \{0, 1, -5, 2, 2, -5, -1, 3, -2, 0, 1, -2, -1, 1, M, 0, 4, 1, -1, -1, 0, -2, -5, 0, -4, 3\},
        /* O */
                   {-2, 0,-4,-1,-1,-4,-3, 2,-2, 0, 3,-3, 0, 0, M, 0, 1, 6, 0,-1, 0,-2, 2, 0,-4, 0},
        /* R */
                   {1,0,0,0,0,0,-3,1,-1,-1,0,0,-3,-2,1,_M,1,-1,0,2,1,0,-1,-2,0,-3,0},
        /* S */
                   { 1, 0,-2, 0, 0,-3, 0,-1, 0, 0, 0,-1,-1, 0,_M, 0,-1,-1, 1, 3, 0, 0,-5, 0,-3, 0},
        /* T */
                    /* U */
 35
                    { 0,-2,-2,-2,-1,-1,-2, 4, 0,-2, 2, 2,-2,_M,-1,-2,-2,-1, 0, 0, 4,-6, 0,-2,-2}
        /* V */
                    {-6,-5,-8,-7,-7, 0,-7,-3,-5, 0,-3,-2,-4,-4,_M,-6,-5, 2,-2,-5, 0,-6,17, 0, 0,-6},
        /* W */
                    /* X */
                    {-3,-3, 0,-4,-4, 7,-5, 0,-1, 0,-4,-1,-2,-2,_M,-5,-4,-4,-3,-3, 0,-2, 0, 0,10,-4},
        /* Y */
                    { 0, 1,-5, 2, 3,-5, 0, 2,-2, 0, 0,-2,-1, 1, M, 0, 3, 0, 0, 0, 0, 0,-2,-6, 0,-4, 4}
 40
        /* Z */
        };
```

45

50

Table 1 (cont')

```
*/
       #include <stdio.h>
 5
       #include <ctype.h>
       #define MAXJMP
                                     16
                                              /* max jumps in a diag */
       #define MAXGAP
                                     24
                                              /* don't continue to penalize gaps larger than this */
       #define JMPS
                                     1024
                                              /* max jmps in an path */
10
       #define MX
                                    4
                                              /* save if there's at least MX-1 bases since last jmp */
                                              /* value of matching bases */
       #define DMAT
                                    3
       #define DMIS
                                    0
                                              /* penalty for mismatched bases */
       #define DINSO
                                    8
                                              /* penalty for a gap */
15
       #define DINS1
                                    1
                                              /* penalty per base */
       #define PINSO
                                              /* penalty for a gap */
                                    8
       #define PINS1
                                              /* penalty per residue */
       struct jmp {
20
                 short
                                    n[MAXJMP];
                                                        /* size of jmp (neg for dely) */
                 unsigned short
                                    x[MAXJMP];
                                                        /* base no. of jmp in seq x */
       };
                                                        /* limits seq to 2^16 -1 */
       struct diag {
25
                 int
                                     score;
                                                        /* score at last jmp */
                 long
                                    offset;
                                                        /* offset of prev block */
                 short
                                                        /* current jmp index */
                                    ijmp;
                 struct jmp
                                                        /* list of jmps */
                                    jp;
       };
30
       struct path {
                 int
                                              /* number of leading spaces */
                 short
                           n[JMPS]; /* size of jmp (gap) */
                           x[JMPS];/* loc of jmp (last elem before gap) */
                 int
35
       };
       char
                           *ofile;
                                                        /* output file name */
                           *namex[2];
       char
                                                        /* seq names: getseqs() */
       char
                           *prog;
                                                        /* prog name for err msgs */
40
       char
                           *seqx[2];
                                                        /* seqs: getseqs() */
                                                        /* best diag: nw() */
                           dmax;
       int
       int
                           dmax0;
                                                        /* final diag */
       int
                           dna:
                                                        /* set if dna: main() */
                                                        /* set if penalizing end gaps */
       int
                           endgaps;
45
       int
                           gapx, gapy;
                                                        /* total gaps in seqs */
                                                        /* seq lens */
       int
                           len0, len1;
       int
                          ngapx, ngapy;
                                                        /* total size of gaps */
       int
                                                        /* max score: nw() */
                           smax;
       int
                           *xbm:
                                                        /* bitmap for matching */
50
       long
                           offset;
                                                        /* current offset in jmp file */
       struct
                 diag
                                                        /* holds diagonals */
                           *dx;
       struct
                 path
                           pp[2];
                                                        /* holds path for segs */
       char
                           *calloc(), *malloc(), *index(), *strcpy();
55
       char
                           *getseq(), *g_calloc();
```

Table 1 (cont')

```
/* Needleman-Wunsch alignment program
        * usage: progs file1 file2
 5
           where file1 and file2 are two dna or two protein sequences.
           The sequences can be in upper- or lower-case an may contain ambiguity
           Any lines beginning with ';', '>' or '<' are ignored
           Max file length is 65535 (limited by unsigned short x in the jmp struct)
           A sequence with 1/3 or more of its elements ACGTU is assumed to be DNA
10
           Output is in the file "align.out"
        * The program may create a tmp file in /tmp to hold info about traceback.
        * Original version developed under BSD 4.3 on a vax 8650
15
       #include "nw.h"
       #include "day.h"
       static
                  _{dbval[26]} = {
                  1,14,2,13,0,0,4,11,0,0,12,0,3,15,0,0,0,5,6,8,8,7,9,0,10,0
20
       };
       static
                  _{pbval[26]} = {
                  1, 2|(1<<('D'-'A'))|(1<<('N'-'A')), 4, 8, 16, 32, 64,
                  128, 256, 0xFFFFFFF, 1<<10, 1<<11, 1<<12, 1<<13, 1<<14,
25
                  1<<15, 1<<16, 1<<17, 1<<18, 1<<19, 1<<20, 1<<21, 1<<22,
                  1<23, 1<24, 1<25[(1<<('E'-'A'))](1<<('Q'-'A'))
       };
       main(ac, av)
                  main
30
                  int
                            ac;
                  char
                            *av[];
        {
                  prog = av[0];
35
                  if (ac!=3) {
                            fprintf(stderr,"usage: %s file1 file2\n", prog);
                            fprintf(stderr,"where file1 and file2 are two dna or two protein sequences.\n"); fprintf(stderr,"The sequences can be in upper- or lower-case\n");
                            fprintf(stderr, "Any lines beginning with ';' or '<' are ignored\n");
40
                            fprintf(stderr,"Output is in the file \"align.out\"\n");
                            exit(1);
                  namex[0] = av[1];
                  namex[1] = av[2];
45
                  seqx[0] = getseq(namex[0], \&len0);
                  seqx[1] = getseq(namex[1], &len1);
                  xbm = (dna)? _dbval : _pbval;
                  endgaps = 0;
                                                           /* 1 to penalize endgaps */
50
                  ofile = "align.out";
                                                 /* output file */
                                      /* fill in the matrix, get the possible jmps */
                  nw();
                  readjmps();
                                      /* get the actual jmps */
                  print();
                                      /* print stats, alignment */
55
                  cleanup(0);
                                      /* unlink any tmp files */
       }
```

```
/* do the alignment, return best score: main()
         * dna: values in Fitch and Smith, PNAS, 80, 1382-1386, 1983
         * pro: PAM 250 values
 5
         * When scores are equal, we prefer mismatches to any gap, prefer
         * a new gap to extending an ongoing gap, and prefer a gap in seqx
         * to a gap in seq y.
         */
        nw()
10
                   nw
        {
                   char
                                         *px, *py;
                                                              /* seqs and ptrs */
                                         *ndely, *dely;
                                                              /* keep track of dely */
                   int
                                                              /* keep track of delx */
                   int
                                         ndelx, delx;
15
                   int
                                         *tmp;
                                                              /* for swapping row0, row1 */
                                                              /* score for each type */
                                         mis;
                   int
                                                              /* insertion penalties */
                   int
                                         ins0, ins1;
                   register
                                         id;
                                                              /* diagonal index */
                   register
                                                              /* jmp index */
                                                              /* score for curr, last row */
20
                   register
                                         *col0, *col1;
                   register
                                                              /* index into segs */
                                         xx, yy;
                   dx = (struct diag *)g_calloc("to get diags", len0+len1+1, sizeof(struct diag));
25
                   ndely = (int *)g_calloc("to get ndely", len1+1, sizeof(int));
                   dely = (int *)g_calloc("to get dely", len1+1, sizeof(int));

col0 = (int *)g_calloc("to get col0", len1+1, sizeof(int));

col1 = (int *)g_calloc("to get col1", len1+1, sizeof(int));
                   ins0 = (dna)? DINS0: PINS0;
30
                   ins1 = (dna)? DINS1: PINS1;
                   smax = -10000;
                   if (endgaps) {
                              for (col0[0] = dely[0] = -ins0, yy = 1; yy \le len1; yy++) {
35
                                         col0[yy] = dely[yy] = col0[yy-1] - ins1;
                                         ndely[yy] = yy;
                              col0[0] = 0;
                                                   /* Waterman Bull Math Biol 84 */
40
                   else
                              for (yy = 1; yy \le len1; yy++)
                                         dely[yy] = -ins0;
                   /* fill in match matrix
45
                   for (px = seqx[0], xx = 1; xx \le len0; px++, xx++) {
                              /* initialize first entry in col
                              if (endgaps) {
50
                                         if(xx == 1)
                                                   col1[0] = delx = -(ins0+ins1);
                                         else
                                                   col1[0] = delx = col0[0] - ins1;
                                         ndelx = xx;
55
                              else {
                                         col1[0] = 0;
                                        delx = -ins0;
                                        ndelx = 0;
60
                             }
```

Table 1 (cont')

```
...nw
                          for (py = seqx[1], yy = 1; yy <= len1; py++, yy++) {
    mis = col0[yy-1];
 5
                                    if (dna)
                                              mis += (xbm[*px-'A']&xbm[*py-'A'])? DMAT : DMIS;
                                     else
                                              mis += _day[*px-'A'][*py-'A'];
10
                                    /* update penalty for del in x seq;
                                     * favor new del over ongong del
                                     * ignore MAXGAP if weighting endgaps
                                    if (endgaps | ndely[yy] < MAXGAP) {
15
                                              if (col0[yy] - ins0 >= dely[yy]) {
                                                        dely[yy] = col0[yy] - (ins0+ins1);
                                                        ndely[yy] = 1;
                                              } else {
                                                        dely[yy] = ins1;
20
                                                        ndely[yy]++;
                                              }
                                    } else {
                                              if (col0[yy] - (ins0+ins1) >= dely[yy]) {
                                                        dely[yy] = col0[yy] - (ins0+ins1);
25
                                                        ndely[yy] = 1;
                                              } else
                                                        ndely[yy]++;
                                    }
                                    /* update penalty for del in y seq;
30
                                     * favor new del over ongong del
                                    if (endgaps | ndelx < MAXGAP) {
                                              if (coll[yy-1] - ins0 >= delx) {
35
                                                        delx = col1[yy-1] - (ins0+ins1);
                                                        ndelx = 1;
                                              } else {
                                                        delx -= ins1;
                                                        ndelx++;
40
                                    } else {
                                              if (col1[yy-1] - (ins0+ins1) >= delx) {
                                                        delx = col1[yy-1] - (ins0+ins1);
                                                        ndelx = 1;
45
                                              } else
                                                        ndelx++;
                                    }
                                    /* pick the maximum score; we're favoring
50
                                     * mis over any del and delx over dely
55
```

```
...nw
                                      id = xx - yy + len1 - 1;
                                       if (mis >= delx && mis >= dely[yy])
 5
                                                 col1[yy] = mis;
                                       else if (delx >= dely[yy]) {
                                                 col1[yy] = delx;
                                                 ij = dx[id].ijmp;
                                                 if (dx[id].jp.n[0] && (!dna || (ndelx >= MAXJMP))
                                                 && xx > dx[id].jp.x[ij]+MX) | mis > dx[id].score+DINS0)) {
10
                                                           dx[id].ijmp++;
                                                           if (++ij >= MAXJMP) {
                                                                      writejmps(id);
                                                                      ij = dx[id].ijmp = 0;
15
                                                                      dx[id].offset = offset;
                                                                      offset += sizeof(struct jmp) + sizeof(offset);
                                                           }
                                                 dx[id].jp.n[ij] = ndelx;
20
                                                 dx[id].jp.x[ij] = xx;
                                                 dx[id].score = delx;
                                       else {
                                                 col1[yy] = dely[yy];
                                                 ij = dx[id].ijmp;
25
                  if (dx[id].jp.n[0] && (!dna || (ndely[yy] >= MAXJMP)
                                                 && xx > dx[id].jp.x[ij]+MX) \parallel mis > dx[id].score+DINS0)) {
                                                           dx[id].ijmp++;
                                                           if (++ij >= MAXJMP) {
30
                                                                      writejmps(id);
                                                                      ij = dx[id].ijmp = 0;
                                                                      dx[id].offset = offset;
                                                                      offset += sizeof(struct jmp) + sizeof(offset);
                                                           }
35
                                                 dx[id].jp.n[ij] = -ndely[yy];
                                                 dx[id].jp.x[ij] = xx;
                                                 dx[id].score = dely[yy];
40
                                       if (xx == len0 && yy < len1) {
                                                 /* last col
                                                 if (endgaps)
                                                           col1[yy] = ins0+ins1*(len1-yy);
45
                                                 if (col1[yy] > smax) {
                                                           smax = col1[yy];
                                                           dmax = id;
                                                 }
                                       }
50
                            if (endgaps && xx < len0)
                                       col1[yy-1] = ins0+ins1*(len0-xx);
                            if (col1[yy-1] > smax) {
                                       smax = col1[yy-1];
55
                                       dmax = id:
                            tmp = col0; col0 = col1; col1 = tmp;
                  (void) free((char *)ndely);
                  (void) free((char *)dely);
(void) free((char *)col0);
(void) free((char *)col1);
60
                                                                     }
```

PCT/US2003/035971

```
* print() -- only routine visible outside this module
 5
        * static:
         * getmat() -- trace back best path, count matches: print()
         * pr_align() -- print alignment of described in array p[]: print()
        * dumpblock() -- dump a block of lines with numbers, stars: pr_align()
        * nums() -- put out a number line: dumpblock()
10
        * putline() -- put out a line (name, [num], seq, [num]): dumpblock()
         * stars() - -put a line of stars: dumpblock()
         * stripname() - strip any path and prefix from a sequame
15
       #include "nw.h"
        #define SPC
                           3
       #define P_LINE
                           256
                                     /* maximum output line */
20
        #define P_SPC
                                     /* space between name or num and seq */
        extern
                 _day[26][26];
       int
                 olen;
                                     /* set output line length */
       FILE
                                     /* output file */
                  *fx;
25
       print()
                 print
        {
                 int
                           lx, ly, firstgap, lastgap;
                                                          /* overlap */
30
                 if ((fx = fopen(ofile, "w")) == 0) {
                           fprintf(stderr,"%s: can't write %s\n", prog, ofile);
                           cleanup(1);
35
                 fprintf(fx, "<first sequence: %s (length = %d)\n", namex[0], len0);
                 fprintf(fx, "<second sequence: %s (length = %d)\n", namex[1], len1);
                 olen = 60;
                 lx = len0;
                 ly = len1;
40
                 firstgap = lastgap = 0;
                 if (dmax < len1 - 1) {
                                                /* leading gap in x */
                           pp[0].spc = firstgap = len1 - dmax - 1;
                           ly = pp[0].spc;
45
                 else if (dmax > len1 - 1) {
                                               /* leading gap in y */
                           pp[1].spc = firstgap = dmax - (len1 - 1);
                           lx = pp[1].spc;
                 if (dmax0 < len0 - 1) {
                                                /* trailing gap in x */
50
                           lastgap = len0 - dmax0 - 1;
                           lx -= lastgap;
                 else if (dmax0 > len0 - 1) { /* trailing gap in y */
                           lastgap = dmax0 - (len0 - 1);
55
                           ly -= lastgap;
                 getmat(lx, ly, firstgap, lastgap);
                 pr_align();
       }
60
```

```
* trace back the best path, count matches
       */
       static
 5
                                                                                                                      getmat
       getmat(lx, ly, firstgap, lastgap)
                                                         /* "core" (minus endgaps) */
                           lx, ly;
                int
                                                         /* leading trailing overlap */
                int
                           firstgap, lastgap;
                                     nm, i0, i1, siz0, siz1;
10
                 int
                                     outx[32];
                 char
                                     pct;
                 double
                                     n0, n1;
                 register
                                     *p0, *p1;
                 register char
15
                 /* get total matches, score
                 i0 = i1 = siz0 = siz1 = 0;
                 p0 = seqx[0] + pp[1].spc;
                 p1 = seqx[1] + pp[0].spc;
20
                 n0 = pp[1].spc + 1;
                 n1 = pp[0].spc + 1;
                  nm = 0;
                  while (*p0 && *p1) {
25
                           if (siz0) {
                                      p1++;
                                      n1++;
                                      siz0--;
30
                            else if (siz1) {
                                      p0++;
                                      n0++;
                                      siz1--;
35
                            else {
                                      if (xbm[*p0-'A']&xbm[*p1-'A'])
                                                 nm++;
                                      if (n0 \mapsto = pp[0].x[i0])
                                                 siz0 = pp[0].n[i0++];
 40
                                      if (n1++ == pp[1].x[i1])
                                                 siz1 = pp[1].n[i1++];
                                      p0++;
                                      p1++;
 45
                            }
                  /* pct homology:
                   * if penalizing endgaps, base is the shorter seq
                   * else, knock off overhangs and take shorter core
 50
                  if (endgaps)
                            lx = (len0 < len1)? len0 : len1;
                             lx = (lx < ly)? lx : ly;
 55
                   pct = 100.*(double)nm/(double)lx;
                  fprintf(fx, "\n");
fprintf(fx, "<%d match%s in an overlap of %d: %.2f percent similarity\n",
                             nm, (nm == 1)? "": "es", lx, pct);
 60
```

```
...getmat
                 fprintf(fx, "<gaps in first sequence: %d", gapx);
                 if (gapx) {
                            (void) sprintf(outx, " (%d %s%s)",
 5
                                      ngapx, (dna)? "base": "residue", (ngapx == 1)? "": "s");
                            fprintf(fx,"%s", outx);
                  fprintf(fx, ", gaps in second sequence: %d", gapy);
10
                  if (gapy) {
                            (void) sprintf(outx, " (%d %s%s)",
                                      ngapy, (dna)? "base": "residue", (ngapy == 1)? "": "s");
                            fprintf(fx,"%s", outx);
                  if (dna)
15
                            fprintf(fx,
                            "\n<score: %d (match = %d, mismatch = %d, gap penalty = %d + %d per base)\n",
                            smax, DMAT, DMIS, DINSO, DINS1);
                  else
20
                             "\n<score: %d (Dayhoff PAM 250 matrix, gap penalty = %d + %d per residue)\n",
                            smax, PINSO, PINS1);
                  if (endgaps)
                            fprintf(fx,
                             "<endgaps penalized. left endgap: %d %s%s, right endgap: %d %s%s\n",
25
                            firstgap, (dna)? "base": "residue", (firstgap == 1)? "": "s", lastgap, (dna)? "base": "residue", (lastgap == 1)? "": "s");
                  else
                             fprintf(fx, "<endgaps not penalized\n");
30
                                                 /* matches in core -- for checking */
         static
                            nm;
                                                 /* lengths of stripped file names */
                            lmax;
         static
                                                 /* jmp index for a path */
         static
                             ij[2];
                                                 /* number at start of current line */
                            nc[2];
         static
                                                 /* current elem number -- for gapping */
                            ni[2];
35
         static
         static
                             siz[2];
                                                 /* ptr to current element */
         static char
                             *ps[2];
                                                  /* ptr to next output char slot */
         static char
                             *po[2];
                            out[2][P_LINE];
                                                 /* output line */
         static char
                             star[P_LINE];
                                                  /* set by stars() */
40
         static char
         * print alignment of described in struct path pp[]
45
        static
                                                                                                                pr_align
        pr_align()
                                                  /* char count */
                   int
                                       nn;
                   int
                                       more;
50
                   register
                   for (i = 0, lmax = 0; i < 2; i++) {
                             nn = stripname(namex[i]);
                             if (nn > lmax)
55
                                       lmax = nn;
                             nc[i] = 1;
                             ni[i] = 1;
                             siz[i] = ij[i] = 0;
 60
                             ps[i] = seqx[i];
                             po[i] = out[i];
                                                                       }
```

```
...pr_align
                 for (nn = nm = 0, more = 1; more;)
                           for (i = more = 0; i < 2; i++) {
 5
                                      * do we have more of this sequence?
                                      */
                                     if (!*ps[i])
                                                continue;
10
                                      more++;
                                     if (pp[i].spc) {
                                                      /* leading space */
                                                *po[i]++='';
15
                                                pp[i].spc--;
                                                         /* in a gap */
                                      else if (siz[i]) {
                                                *po[i]++='-';
                                                siz[i]--;
20
                                      else {
                                                         /* we're putting a seq element
                                                *po[i] = *ps[i];
                                                if (islower(*ps[i]))
25
                                                         *ps[i] = toupper(*ps[i]);
                                                po[i]++;
                                                ps[i]++;
30
                                                * are we at next gap for this seq?
                                                */
                                                if (ni[i] == pp[i].x[ij[i]]) \{
                                                          * we need to merge all gaps
35
                                                          * at this location
                                                         siz[i] = pp[i].n[ij[i]++];
                                                          while (ni[i] = pp[i].x[ij[i]])
                                                                   siz[i] \leftarrow pp[i].n[ij[i]++];
40
                                                ni[i]++;
                                      }
                           if (++nn == olen | !more && nn) {
45
                                      dumpblock();
                                      for (i = 0; i < 2; i++)
                                               po[i] = out[i];
                                      nn = 0;
                           }
50
                 }
       }
        * dump a block of lines, including numbers, stars: pr_align()
55
       static
       dumpblock()
                 dumpblock
       {
60
                 register i;
                 for (i = 0; i < 2; i++)
                           po[i] - = '0';
```

Table 1 (cont')

...dumpblock (void) putc('\n', fx); 5 for (i = 0; i < 2; i++) { if (*out[i] && (*out[i] != ' ' || *(po[i]) != ' ')) { if (i == 0)nums(i); **if** (i = 0 && *out[1]) 10 stars(); putline(i); if (i = 0 & *out[1])fprintf(fx, star); if (i == 1)15 nums(i); } } } 20 * put out a number line: dumpblock() */ static nums nums(ix) 25 /* index in out[] holding seq line */ int ix; nline[P_LINE]; char register i, j; register char *pn, *px, *py; 30 for $(pn = nline, i = 0; i < lmax+P_SPC; i++, pn++)$ *pn = ' '; for $(i = nc[ix], py = out[ix]; *py; py++, pn++) {$ if (*py == ' ' || *py == '-') 35 *pn = ' '; else { if (i%10 = 0 || (i = 1 && nc[ix] != 1)) { j = (i < 0)? -i : i;for (px = pn; j; j /= 10, px-)40 px = j%10 + '0';if (i < 0)*px = '-'; else 45 *pn = ' '; i++; } *pn = '0';50 nc[ix] = i;for (pn = nline; *pn; pn++)(void) putc(*pn, fx); (void) putc('\n', fx); } 55 * put out a line (name, [num], seq, [num]): dumpblock() static

{

60

putline(ix)

int

ix;

putline

Table 1 (cont')

...putline

```
int
                 register char
 5
                                      *px;
                 for (px = namex[ix], i = 0; *px && *px != ':'; px++, i++)
                 (void) putc(*px, fx);
for (; i < lmax+P_SPC; i++)
10
                           (void) putc('', fx);
                 /* these count from 1:
                  * ni[] is current element (from 1)
                  * nc[] is number at start of current line
15
                  for (px = out[ix]; *px; px++)
                            (void) putc(*px&0x7F, fx);
                  (void) putc('\n', fx);
       }
20
         * put a line of stars (seqs always in out[0], out[1]): dumpblock()
        static
25
        stars()
                  stars
        {
                                       *p0, *p1, cx, *px;
                  register char
30
                  if (!*out[0] || (*out[0] == '' && *(po[0]) == '') ||
                     !*out[1] || (*out[1] = '' && *(po[1]) == '"))
                  px = star;
35
                  for (i = lmax+P\_SPC; i; i-)
                             *px++='';
                  for (p0 = out[0], p1 = out[1]; *p0 && *p1; p0++, p1++) {
                            if (isalpha(*p0) && isalpha(*p1)) {
 40
                                       if (xbm[*p0-'A']&xbm[*p1-'A']) {
                                                 cx = '*';
                                                 nm++;
 45
                                       else if (!dna && _day[*p0-'A'][*p1-'A'] > 0)
                                                 cx = '.';
                                       else
                                                 cx = ' ';
 50
                             }
                             else
                                       cx = ' ';
                             *px++=cx;
                   *px++ = 'n';
 55
                   *px = '0';
         }
```

60

```
* strip path or prefix from pn, return len: pr_align()
 5
       static
       stripname(pn)
                  stripname
                                       /* file name (may be path) */
                  char
                            *pn;
                  register char
                                       *px, *py;
10
                 py = 0;
for (px = pn; *px; px++)
if (*px == '/')
py =
                                       py = px + 1;
15
                  if (py)
                  (void) strcpy(pn, py);
return(strlen(pn));
20
        }
25
 30
 35
 40
 45
 50
  55
  60
```

```
* cleanup() -- cleanup any tmp file
         * getseq() -- read in seq, set dna, len, maxlen
 5
         * g_calloc() -- calloc() with error checkin
         * readjmps() -- get the good jmps, from tmp file if necessary
         * writejmps() -- write a filled array of jmps to a tmp file: nw()
         */
        #include "nw.h"
10
        #include <sys/file.h>
                   *jname = "/tmp/homgXXXXXX";
        char
                                                                          /* tmp file for jmps */
        FILE
15
        int
                   cleanup();
                                                                          /* cleanup tmp file */
        long
                   lseek();
         * remove any tmp file if we blow
20
         */
                                                                                                                                 cleanup
        cleanup(i)
                   int
                              i;
        {
                   if (fj)
25
                              (void) unlink(jname);
                   exit(i);
        }
         * read, return ptr to seq, set dna, len, maxlen * skip lines starting with ';', '<', or '>'
30
         * seq in upper or lower case
         */
        char
35
        getseq(file, len)
                                                                                                                                 getseq
                              *file;
                                         /* file name */
                   char
                   int
                              *len;
                                        /* seq len */
        {
                   char
                                         line[1024], *pseq;
40
                   register char
                                         *px, *py;
                   int
                                         natgc, tlen;
                   FILE
                                         *fp;
                   if ((fp = fopen(file,"r")) == 0) {
45
                              fprintf(stderr,"%s: can't read %s\n", prog, file);
                              exit(1);
                   tlen = natgc = 0;
                   while (fgets(line, 1024, fp)) {
    if (*line == ';' || *line == '<' || *line == '>')
50
                                        continue;
                              for (px = line; *px != '\n'; px++)
if (isupper(*px) || islower(*px))
                                                   tlen++;
55
                   if ((pseq = malloc((unsigned)(tlen+6))) == 0) {
                              fprintf(stderr,"%s: malloc() failed to get %d bytes for %s\n", prog, tlen+6, file);
60
                   pseq[0] = pseq[1] = pseq[2] = pseq[3] = \0;
```

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```
...getseq
                  py = pseq + 4;
                  *len = tlen;
 5
                  rewind(fp);
                  while (fgets(line, 1024, fp)) {
    if (*line == ';' || *line == '<' || *line == '>')
                                      continue;
10
                            for (px = line; *px != 'n'; px++) {
                                      if (isupper(*px))
                                                 *py++ = *px;
                                      else if (islower(*px))
                                      *py++ = toupper(*px);
if (index("ATGCU",*(py-1)))
15
                                                 natgc++;
                            }
                  *py++ = '\0';
20
                  *py = '0';
                  (void) fclose(fp);
                  dna = natgc > (tlen/3);
                  return(pseq+4);
        }
25
        char
                                                                                                                          g_calloc
        g_calloc(msg, nx, sz)
                                                 /* program, calling routine */
                  char
                            *msg;
                                                /* number and size of elements */
                  int
                            nx, sz;
30
        {
                                      *px, *calloc();
                  char
                  if ((px = calloc((unsigned)nx, (unsigned)sz)) == 0) {
                            if (*msg) {
                                      fprintf(stderr, "%s: g_calloc() failed %s (n=%d, sz=%d)\n", prog, msg, nx, sz);
35
                  return(px);
40
        }
        * get final jmps from dx[] or tmp file, set pp[], reset dmax: main()
45
        readjmps()
                  readjmps
        {
                  int
                                      fd = -1;
                                      siz, i0, i1;
                  int
50
                  register i, j, xx;
                  if (fj) {
                            (void) fclose(fj);
                            if ((fd = open(jname, O_RDONLY, 0)) < 0) {
55
                                       fprintf(stderr, "%s: can't open() %s\n", prog, jname);
                                       cleanup(1);
                            }
                  for (i = i0 = i1 = 0, dmax0 = dmax, xx = len0; ; i++) {
60
                            while (1) {
                                      for (j = dx[dmax].ijmp; j \ge 0 && dx[dmax].jp.x[j] \ge xx; j-)
```

Table 1 (cont')

...readjmps

```
if (j < 0 && dx[dmax].offset && fj) {
                                                 (void) lseek(fd, dx[dmax].offset, 0);
 5
                                                 (void) read(fd, (char *)&dx[dmax].jp, sizeof(struct jmp));
                                                 (void) read(fd, (char *)&dx[dmax].offset, sizeof(dx[dmax].offset));
                                                 dx[dmax].ijmp = MAXJMP-1;
                                       else
10
                                                 break;
                            if (i >= JMPS) {
                                       fprintf(stderr, "%s: too many gaps in alignment\n", prog);
                                       cleanup(1);
                            if (j >= 0) {
15
                                       siz = dx[dmax].jp.n[j];
                                       xx = dx[dmax].jp.x[j];
                                       dmax += siz;
20
                                      if (siz < 0) {
                                                                      /* gap in second seq */
                                                 pp[1].n[i1] = -siz;
                                                 xx += siz;
                                                 /* id = xx - yy + len1 - 1
                                                  */
25
                                                 pp[1].x[i1] = xx - dmax + len1 - 1;
                                                 gapy++;
                                                 ngapy -= siz;
        /* ignore MAXGAP when doing endgaps */
                                                 siz = (-siz < MAXGAP || endgaps)? -siz : MAXGAP;
30
                                                 i1++;
                                       else if (siz > 0) { /* gap in first seq */
                                                 pp[0].n[i0] = siz;
                                                 pp[0].x[i0] = xx;
35
                                                 gapx++;
                                                 ngapx += siz;
       /* ignore MAXGAP when doing endgaps */
                                                 siz = (siz < MAXGAP || endgaps)? siz : MAXGAP;
                                                 i0++;
40
                                      }
                            else
                                       break;
                  }
45
                  /* reverse the order of jmps
                  for (j = 0, i0--; j < i0; j++, i0--)
                            i = pp[0].n[j]; pp[0].n[j] = pp[0].n[i0]; pp[0].n[i0] = i; i = pp[0].x[j]; pp[0].x[j] = pp[0].x[i0]; pp[0].x[i0] = i;
50
                  for (j = 0, i1-; j < i1; j++, i1-)
                            i = pp[1].n[j]; pp[1].n[j] = pp[1].n[i1]; pp[1].n[i1] = i;
                            i = pp[1].x[j]; pp[1].x[j] = pp[1].x[i1]; pp[1].x[i1] = i;
55
                  if (fd >= 0)
                            (void) close(fd);
                  if (fj) {
                            (void) unlink(jname);
60
                            fj = 0;
                            offset = 0;
                 }
                                                           }
```

```
* write a filled jmp struct offset of the prev one (if any): nw()
 5
         writejmps(ix)
                      writejmps
         {
10
                      char
                                    *mktemp();
                      if (!fj) {
                                    if (mktemp(jname) < 0) {
    fprintf(stderr, "%s: can't mktemp() %s\n", prog, jname);</pre>
                                                 cleanup(1);
15
                                   if ((fj = fopen(jname, "w")) == 0) {
    fprintf(stderr, "%s: can't write %s\n", prog, jname);
20
                                    }
                       (void) fwrite((char *)&dx[ix].jp, sizeof(struct jmp), 1, fj);
(void) fwrite((char *)&dx[ix].offset, sizeof(dx[ix].offset), 1, fj);
25
```

Table 2

5 PRO XXXXXXXXXXXXXXX (Length = 15 amino acids)

Comparison Protein XXXXXYYYYYYYY (Length = 12 amino acids)

% amino acid sequence identity =

(the number of identically matching amino acid residues between the two polypeptide sequences as

determined by ALIGN-2) divided by (the total number of amino acid residues of the PRO polypeptide) =

5 divided by 15 = 33.3%

Table 3

15 PRO XXXXXXXXXX (Length = 10 amino acids)

Comparison Protein XXXXXYYYYYYZZYZ (Length = 15 amino acids)

% amino acid sequence identity =

(the number of identically matching amino acid residues between the two polypeptide sequences as

determined by ALIGN-2) divided by (the total number of amino acid residues of the PRO polypeptide) =

5 divided by 10 = 50%

Table 4

25 PRO-DNA NNNNNNNNNNNNN (Length = 14 nucleotides)

Comparison DNA NNNNNNLLLLLLLLLL (Length = 16 nucleotides)

% nucleic acid sequence identity =

30 (the number of identically matching nucleotides between the two nucleic acid sequences as determined by ALIGN-2) divided by (the total number of nucleotides of the PRO-DNA nucleic acid sequence) = 6 divided by 14 = 42.9%

Table 5

35

PRO-DNA NNNNNNNNNN (Length = 12 nucleotides)

Comparison DNA NNNNLLLVV (Length = 9 nucleotides)

% nucleic acid sequence identity =

40

(the number of identically matching nucleotides between the two nucleic acid sequences as determined by ALIGN-2) divided by (the total number of nucleotides of the PRO-DNA nucleic acid sequence) = 4 divided by 12 = 33.3%

II. Compositions and Methods of the Invention

A. Full-Length PRO Polypeptides

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The present invention provides newly identified and isolated nucleotide sequences encoding polypeptides referred to in the present application as PRO polypeptides. In particular, cDNAs encoding various PRO polypeptides have been identified and isolated, as disclosed in further detail in the Examples below. However, for sake of simplicity, in the present specification the protein encoded by the full length native nucleic acid molecules disclosed herein as well as all further native homologues and variants included in the foregoing definition of PRO, will be referred to as "PRO/number", regardless of their origin or mode of preparation.

As disclosed in the Examples below, various cDNA clones have been disclosed. The predicted amino acid sequence can be determined from the nucleotide sequence using routine skill. For the PRO polypeptides and encoding nucleic acids described herein, Applicants have identified what is believed to be the reading frame best identifiable with the sequence information available at the time.

B. PRO Polypeptide Variants

In addition to the full-length native sequence PRO polypeptides described herein, it is contemplated that PRO variants can be prepared. PRO variants can be prepared by introducing appropriate nucleotide changes into the PRO DNA, and/or by synthesis of the desired PRO polypeptide. Those skilled in the art will appreciate that amino acid changes may alter post-translational processes of the PRO, such as changing the number or position of glycosylation sites or altering the membrane anchoring characteristics.

Variations in the native full-length sequence PRO or in various domains of the PRO described herein, can be made, for example, using any of the techniques and guidelines for conservative and non-conservative mutations set forth, for instance, in U.S. Patent No. 5,364,934. Variations may be a substitution, deletion or insertion of one or more codons encoding the PRO that results in a change in the amino acid sequence of the PRO as compared with the native sequence PRO. Optionally, the variation is by substitution of at least one amino acid with any other amino acid in one or more of the domains of the PRO. Guidance in determining which amino acid residue may be inserted, substituted or deleted without adversely affecting the desired activity may be found by comparing the sequence of the PRO with that of homologous known protein molecules and minimizing the number of amino acid sequence changes made in regions of high homology. Amino acid substitutions can be the result of replacing one amino acid with another amino acid having similar structural and/or chemical properties, such as the replacement of a leucine with a serine, i.e., conservative amino acid replacements. Insertions or deletions may optionally be in the range of about 1 to 5 amino acids. The variation allowed may be determined by systematically making insertions, deletions or substitutions of amino acids in the sequence and testing the resulting variants for activity exhibited by the full-length or mature native sequence.

PRO polypeptide fragments are provided herein. Such fragments may be truncated at the N-terminus or C-terminus, or may lack internal residues, for example, when compared with a full length native

protein. Certain fragments lack amino acid residues that are not essential for a desired biological activity of the PRO polypeptide.

PRO fragments may be prepared by any of a number of conventional techniques. Desired peptide fragments may be chemically synthesized. An alternative approach involves generating PRO fragments by enzymatic digestion, e.g., by treating the protein with an enzyme known to cleave proteins at sites defined by particular amino acid residues, or by digesting the DNA with suitable restriction enzymes and isolating the desired fragment. Yet another suitable technique involves isolating and amplifying a DNA fragment encoding a desired polypeptide fragment, by polymerase chain reaction (PCR). Oligonucleotides that define the desired termini of the DNA fragment are employed at the 5' and 3' primers in the PCR. Preferably, PRO polypeptide fragments share at least one biological and/or immunological activity with the native PRO polypeptide disclosed herein.

In particular embodiments, conservative substitutions of interest are shown in Table 6 under the heading of preferred substitutions. If such substitutions result in a change in biological activity, then more substantial changes, denominated exemplary substitutions in Table 6, or as further described below in reference to amino acid classes, are introduced and the products screened.

Table 6

| | Original | Exemplary | Preferred |
|----|----------|--------------------------|---------------|
| | Residue | Substitutions | Substitutions |
| 20 | Ala (A) | val; leu; ile | val |
| | Arg (R) | lys; gln; asn | lys |
| | Asn (N) | gln; his; lys; arg | gln |
| | Asp (D) | glu | glu |
| | Cys (C) | ser | ser |
| | Gln (Q) | asn | asn |
| | Glu (E) | asp | asp |
| | Gly (G) | pro; ala | ala |
| 30 | His (H) | asn; gln; lys; arg | arg |
| | Ile (I) | leu; val; met; ala; phe; | |
| | | norleucine | leu |
| | Leu (L) | norleucine; ile; val; | |
| | | met; ala; phe | ile |
| 35 | Lys (K) | arg; gln; asn | arg |
| | Met (M) | leu; phe; ile | leu |
| | Phe (F) | leu; val; ile; ala; tyr | leu |
| | Pro (P) | ala | ala |
| | Ser (S) | thr | thr |
| 40 | Thr (T) | ser | ser |
| | Trp (W) | tyr; phe | tyr |
| | Tyr (Y) | trp; phe; thr; ser | phe |
| | Val (V) | ile; leu; met; phe; | |
| | | ala; norleucine | leu |

Substantial modifications in function or immunological identity of the PRO polypeptide are accomplished by selecting substitutions that differ significantly in their effect on maintaining (a) the structure of the polypeptide backbone in the area of the substitution, for example, as a sheet or helical conformation, (b) the charge or hydrophobicity of the molecule at the target site, or (c) the bulk of the side chain. Naturally occurring residues are divided into groups based on common side-chain properties:

(1) hydrophobic: norleucine, met, ala, val, leu, ile;

5

10

15

45

- (2) neutral hydrophilic: cys, ser, thr;
- (3) acidic: asp, glu;
- (4) basic: asn, gln, his, lys, arg;
- (5) residues that influence chain orientation: gly, pro; and
- (6) aromatic: trp, tyr, phe.

5

10

15

20

25

30

35

Non-conservative substitutions will entail exchanging a member of one of these classes for another class. Such substituted residues also may be introduced into the conservative substitution sites or, more preferably, into the remaining (non-conserved) sites.

The variations can be made using methods known in the art such as oligonucleotide-mediated (site-directed) mutagenesis, alanine scanning, and PCR mutagenesis. Site-directed mutagenesis [Carter et al., Nucl. Acids Res., 13:4331 (1986); Zoller et al., Nucl. Acids Res., 10:6487 (1987)], cassette mutagenesis [Wells et al., Gene, 34:315 (1985)], restriction selection mutagenesis [Wells et al., Philos. Trans. R. Soc. London SerA, 317:415 (1986)] or other known techniques can be performed on the cloned DNA to produce the PRO variant DNA.

Scanning amino acid analysis can also be employed to identify one or more amino acids along a contiguous sequence. Among the preferred scanning amino acids are relatively small, neutral amino acids. Such amino acids include alanine, glycine, serine, and cysteine. Alanine is typically a preferred scanning amino acid among this group because it eliminates the side-chain beyond the beta-carbon and is less likely to alter the main-chain conformation of the variant [Cunningham and Wells, Science, 244: 1081-1085 (1989)]. Alanine is also typically preferred because it is the most common amino acid. Further, it is frequently found in both buried and exposed positions [Creighton, The Proteins, (W.H. Freeman & Co., N.Y.); Chothia, J. Mol. Biol., 150:1 (1976)]. If alanine substitution does not yield adequate amounts of variant, an isoteric amino acid can be used.

C. Modifications of PRO

Covalent modifications of PRO are included within the scope of this invention. One type of covalent modification includes reacting targeted amino acid residues of a PRO polypeptide with an organic derivatizing agent that is capable of reacting with selected side chains or the N- or C- terminal residues of the PRO. Derivatization with bifunctional agents is useful, for instance, for crosslinking PRO to a water-insoluble support matrix or surface for use in the method for purifying anti-PRO antibodies, and vice-versa. Commonly used crosslinking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxysuccinimide esters, for example, esters with 4-azidosalicylic acid, homobifunctional imidoesters, including disuccinimidyl esters such as 3,3'-dithiobis(succinimidylpropionate), bifunctional maleimides such as bis-N-maleimido-1,8-octane and agents such as methyl-3-[(p-azidophenyl)dithio]propioimidate.

Other modifications include deamidation of glutaminyl and asparaginyl residues to the corresponding glutamyl and aspartyl residues, respectively, hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl or threonyl residues, methylation of the α -amino groups of lysine, arginine, and histidine side chains [T.E. Creighton, <u>Proteins: Structure and Molecular Properties</u>, W.H. Freeman & Co., San Francisco, pp. 79-86 (1983)], acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl group.

Another type of covalent modification of the PRO polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. "Altering the native glycosylation pattern" is intended for purposes herein to mean deleting one or more carbohydrate moieties found in native sequence PRO (either by removing the underlying glycosylation site or by deleting the glycosylation by chemical and/or enzymatic means), and/or adding one or more glycosylation sites that are not present in the native sequence PRO. In addition, the phrase includes qualitative changes in the glycosylation of the native proteins, involving a change in the nature and proportions of the various carbohydrate moieties present.

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Addition of glycosylation sites to the PRO polypeptide may be accomplished by altering the amino acid sequence. The alteration may be made, for example, by the addition of, or substitution by, one or more serine or threonine residues to the native sequence PRO (for O-linked glycosylation sites). The PRO amino acid sequence may optionally be altered through changes at the DNA level, particularly by mutating the DNA encoding the PRO polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids.

Another means of increasing the number of carbohydrate moieties on the PRO polypeptide is by chemical or enzymatic coupling of glycosides to the polypeptide. Such methods are described in the art, e.g., in WO 87/05330 published 11 September 1987, and in Aplin and Wriston, <u>CRC Crit. Rev. Biochem.</u>, pp. 259-306 (1981).

Removal of carbohydrate moieties present on the PRO polypeptide may be accomplished chemically or enzymatically or by mutational substitution of codons encoding for amino acid residues that serve as targets for glycosylation. Chemical deglycosylation techniques are known in the art and described, for instance, by Hakimuddin, et al., Arch. Biochem. Biophys., 259:52 (1987) and by Edge et al., Anal. Biochem., 118:131 (1981). Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of a variety of endo- and exo-glycosidases as described by Thotakura et al., Meth. Enzymol., 138:350 (1987).

Another type of covalent modification of PRO comprises linking the PRO polypeptide to one of a variety of nonproteinaceous polymers, e.g., polyethylene glycol (PEG), polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,144; 4,670,417; 4,791,192 or 4,179,337.

The PRO of the present invention may also be modified in a way to form a chimeric molecule comprising PRO fused to another, heterologous polypeptide or amino acid sequence.

In one embodiment, such a chimeric molecule comprises a fusion of the PRO with a tag polypeptide which provides an epitope to which an anti-tag antibody can selectively bind. The epitope tag is generally placed at the amino- or carboxyl- terminus of the PRO. The presence of such epitope-tagged forms of the PRO can be detected using an antibody against the tag polypeptide. Also, provision of the epitope tag enables the PRO to be readily purified by affinity purification using an anti-tag antibody or another type of affinity matrix that binds to the epitope tag. Various tag polypeptides and their respective antibodies are well known in the art. Examples include poly-histidine (poly-his) or poly-histidine-glycine (poly-his-gly) tags; the flu HA tag polypeptide and its antibody 12CA5 [Field et al., Mol. Cell. Biol., 8:2159-2165 (1988)]; the c-myc tag and the 8F9, 3C7, 6E10, G4, B7 and 9E10 antibodies thereto [Evan et al.,

Molecular and Cellular Biology, 5:3610-3616 (1985)]; and the Herpes Simplex virus glycoprotein D (gD) tag and its antibody [Paborsky et al., Protein Engineering, 3(6):547-553 (1990)]. Other tag polypeptides include the Flag-peptide [Hopp et al., BioTechnology, 6:1204-1210 (1988)]; the KT3 epitope peptide [Martin et al., Science, 255:192-194 (1992)]; an alpha-tubulin epitope peptide [Skinner et al., J. Biol. Chem., 266:15163-15166 (1991)]; and the T7 gene 10 protein peptide tag [Lutz-Freyermuth et al., Proc. Natl. Acad. Sci. USA, 87:6393-6397 (1990)].

In an alternative embodiment, the chimeric molecule may comprise a fusion of the PRO with an immunoglobulin or a particular region of an immunoglobulin. For a bivalent form of the chimeric molecule (also referred to as an "immunoadhesin"), such a fusion could be to the Fc region of an IgG molecule. The Ig fusions preferably include the substitution of a soluble (transmembrane domain deleted or inactivated) form of a PRO polypeptide in place of at least one variable region within an Ig molecule. In a particularly preferred embodiment, the immunoglobulin fusion includes the hinge, CH2 and CH3, or the hinge, CH1, CH2 and CH3 regions of an IgG1 molecule. For the production of immunoglobulin fusions see also US Patent No. 5,428,130 issued June 27, 1995.

D. Preparation of PRO

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The description below relates primarily to production of PRO by culturing cells transformed or transfected with a vector containing PRO nucleic acid. It is, of course, contemplated that alternative methods, which are well known in the art, may be employed to prepare PRO. For instance, the PRO sequence, or portions thereof, may be produced by direct peptide synthesis using solid-phase techniques [see, e.g., Stewart et al., Solid-Phase Peptide Synthesis, W.H. Freeman Co., San Francisco, CA (1969); Merrifield, J. Am. Chem. Soc., 85:2149-2154 (1963)]. In vitro protein synthesis may be performed using manual techniques or by automation. Automated synthesis may be accomplished, for instance, using an Applied Biosystems Peptide Synthesizer (Foster City, CA) using manufacturer's instructions. Various portions of the PRO may be chemically synthesized separately and combined using chemical or enzymatic methods to produce the full-length PRO.

1. Isolation of DNA Encoding PRO

DNA encoding PRO may be obtained from a cDNA library prepared from tissue believed to possess the PRO mRNA and to express it at a detectable level. Accordingly, human PRO DNA can be conveniently obtained from a cDNA library prepared from human tissue, such as described in the Examples. The PRO-encoding gene may also be obtained from a genomic library or by known synthetic procedures (e.g., automated nucleic acid synthesis).

Libraries can be screened with probes (such as antibodies to the PRO or oligonucleotides of at least about 20-80 bases) designed to identify the gene of interest or the protein encoded by it. Screening the cDNA or genomic library with the selected probe may be conducted using standard procedures, such as described in Sambrook et al., Molecular Cloning: A Laboratory Manual (New York: Cold Spring Harbor Laboratory Press, 1989). An alternative means to isolate the gene encoding PRO is to use PCR methodology [Sambrook et al., supra; Dieffenbach et al., PCR Primer: A Laboratory Manual (Cold Spring Harbor Laboratory Press, 1995)].

The Examples below describe techniques for screening a cDNA library. The oligonucleotide sequences selected as probes should be of sufficient length and sufficiently unambiguous that false positives

are minimized. The oligonucleotide is preferably labeled such that it can be detected upon hybridization to DNA in the library being screened. Methods of labeling are well known in the art, and include the use of radiolabels like ³²P-labeled ATP, biotinylation or enzyme labeling. Hybridization conditions, including moderate stringency and high stringency, are provided in Sambrook et al., <u>supra</u>.

Sequences identified in such library screening methods can be compared and aligned to other known sequences deposited and available in public databases such as GenBank or other private sequence databases. Sequence identity (at either the amino acid or nucleotide level) within defined regions of the molecule or across the full-length sequence can be determined using methods known in the art and as described herein.

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Nucleic acid having protein coding sequence may be obtained by screening selected cDNA or genomic libraries using the deduced amino acid sequence disclosed herein for the first time, and, if necessary, using conventional primer extension procedures as described in Sambrook et al., supra, to detect precursors and processing intermediates of mRNA that may not have been reverse-transcribed into cDNA.

2. Selection and Transformation of Host Cells

Host cells are transfected or transformed with expression or cloning vectors described herein for PRO production and cultured in conventional nutrient media modified as appropriate for inducing promoters, selecting transformants, or amplifying the genes encoding the desired sequences. The culture conditions, such as media, temperature, pH and the like, can be selected by the skilled artisan without undue experimentation. In general, principles, protocols, and practical techniques for maximizing the productivity of cell cultures can be found in Mammalian Cell Biotechnology: a Practical Approach, M. Butler, ed. (IRL Press, 1991) and Sambrook et al., supra.

Methods of eukaryotic cell transfection and prokaryotic cell transformation are known to the ordinarily skilled artisan, for example, CaCl₂, CaPO₄, liposome-mediated and electroporation. Depending on the host cell used, transformation is performed using standard techniques appropriate to such cells. The calcium treatment employing calcium chloride, as described in Sambrook et al., supra, or electroporation is generally used for prokaryotes. Infection with Agrobacterium tumefaciens is used for transformation of certain plant cells, as described by Shaw et al., Gene, 23:315 (1983) and WO 89/05859 published 29 June 1989. For mammalian cells without such cell walls, the calcium phosphate precipitation method of Graham and van der Eb, Virology, 52:456-457 (1978) can be employed. General aspects of mammalian cell host system transfections have been described in U.S. Patent No. 4,399,216. Transformations into yeast are typically carried out according to the method of Van Solingen et al., J. Bact., 130:946 (1977) and Hsiao et al., Proc. Natl. Acad. Sci. (USA), 76:3829 (1979). However, other methods for introducing DNA into cells, such as by nuclear microinjection, electroporation, bacterial protoplast fusion with intact cells, or polycations, e.g., polybrene, polyornithine, may also be used. For various techniques for transforming mammalian cells, see Keown et al., Methods in Enzymology, 185:527-537 (1990) and Mansour et al., Nature, 336:348-352 (1988).

Suitable host cells for cloning or expressing the DNA in the vectors herein include prokaryote, yeast, or higher eukaryote cells. Suitable prokaryotes include but are not limited to eubacteria, such as Gram-negative or Gram-positive organisms, for example, Enterobacteriaceae such as *E. coli*. Various *E. coli*

strains are publicly available, such as E. coli K12 strain MM294 (ATCC 31,446); E. coli X1776 (ATCC 31,537); E. coli strain W3110 (ATCC 27,325) and K5 772 (ATCC 53,635). Other suitable prokaryotic host cells include Enterobacteriaceae such as Escherichia, e.g., E. coli, Enterobacter, Erwinia, Klebsiella, Proteus, Salmonella, e.g., Salmonella typhimurium, Serratia, e.g., Serratia marcescans, and Shigella, as well as Bacilli such as B. subtilis and B. licheniformis (e.g., B. licheniformis 41P disclosed in DD 266,710 published 12 April 1989), Pseudomonas such as P. aeruginosa, and Streptomyces. These examples are illustrative rather than limiting. Strain W3110 is one particularly preferred host or parent host because it is a common host strain for recombinant DNA product fermentations. Preferably, the host cell secretes minimal amounts of proteolytic enzymes. For example, strain W3110 may be modified to effect a genetic mutation in the genes encoding proteins endogenous to the host, with examples of such hosts including E. coli W3110 strain 1A2, which has the complete genotype tonA; E. coli W3110 strain 9E4, which has the complete genotype tonA ptr3; E. coli W3110 strain 27C7 (ATCC 55,244), which has the complete genotype tonA ptr3 phoA E15 (argF-lac)169 degP ompT kanr; E. coli W3110 strain 37D6, which has the complete genotype tonA ptr3 phoA E15 (argF-lac)169 degP ompT rbs7 ilvG kan'; E. coli W3110 strain 40B4, which is strain 37D6 with a non-kanamycin resistant degP deletion mutation; and an E. coli strain having mutant periplasmic protease disclosed in U.S. Patent No. 4,946,783 issued 7 August 1990. Alternatively, in vitro methods of cloning, e.g., PCR or other nucleic acid polymerase reactions, are suitable.

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In addition to prokaryotes, eukaryotic microbes such as filamentous fungi or yeast are suitable cloning or expression hosts for PRO-encoding vectors. Saccharomyces cerevisiae is a commonly used lower eukaryotic host microorganism. Others include Schizosaccharomyces pombe (Beach and Nurse, Nature, 290: 140 [1981]; EP 139,383 published 2 May 1985); Kluyveromyces hosts (U.S. Patent No. 4,943,529; Fleer et al., Bio/Technology, 9:968-975 (1991)) such as, e.g., K. lactis (MW98-8C, CBS683, CBS4574; Louvencourt et al., J. Bacteriol., 154(2):737-742 [1983]), K. fragilis (ATCC 12,424), K. bulgaricus (ATCC 16,045), K. wickeramii (ATCC 24,178), K. waltii (ATCC 56,500), K. drosophilarum (ATCC 36,906; Van den Berg et al., Bio/Technology, 8:135 (1990)), K. thermotolerans, and K. marxianus; yarrowia (EP 402,226); Pichia pastoris (EP 183,070; Sreekrishna et al., J. Basic Microbiol., 28:265-278 [1988]); Candida; Trichoderma reesia (EP 244,234); Neurospora crassa (Case et al., Proc. Natl. Acad. Sci. USA, 76:5259-5263 [1979]); Schwanniomyces such as Schwanniomyces occidentalis (EP 394,538 published 31 October 1990); and filamentous fungi such as, e.g., Neurospora, Penicillium, Tolypocladium (WO 91/00357 published 10 January 1991), and Aspergillus hosts such as A. nidulans (Ballance et al., Biochem. Biophys. Res. Commun., 112:284-289 [1983]; Tilburn et al., Gene, 26:205-221 [1983]; Yelton et al., Proc. Natl. Acad. Sci. USA, 81: 1470-1474 [1984]) and A. niger (Kelly and Hynes, EMBO J., 4:475-479 [1985]). Methylotropic yeasts are suitable herein and include, but are not limited to, yeast capable of growth on methanol selected from the genera consisting of Hansenula, Candida, Kloeckera, Pichia, Saccharomyces, Torulopsis, and Rhodotorula. A list of specific species that are exemplary of this class of yeasts may be found in C. Anthony, The Biochemistry of Methylotrophs, 269 (1982).

Suitable host cells for the expression of glycosylated PRO are derived from multicellular organisms. Examples of invertebrate cells include insect cells such as Drosophila S2 and Spodoptera Sf9, as well as plant cells. Examples of useful mammalian host cell lines include Chinese hamster ovary (CHO) and COS cells. More specific examples include monkey kidney CV1 line transformed by SV40 (COS-7, ATCC

CRL 1651); human embryonic kidney line (293 or 293 cells subcloned for growth in suspension culture, Graham et al., J. Gen Virol., 36:59 (1977)); Chinese hamster ovary cells/-DHFR (CHO, Urlaub and Chasin, Proc. Natl. Acad. Sci. USA, 77:4216 (1980)); mouse sertoli cells (TM4, Mather, Biol. Reprod., 23:243-251 (1980)); human lung cells (W138, ATCC CCL 75); human liver cells (Hep G2, HB 8065); and mouse mammary tumor (MMT 060562, ATCC CCL51). The selection of the appropriate host cell is deemed to be within the skill in the art.

3. Selection and Use of a Replicable Vector

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The nucleic acid (e.g., cDNA or genomic DNA) encoding PRO may be inserted into a replicable vector for cloning (amplification of the DNA) or for expression. Various vectors are publicly available. The vector may, for example, be in the form of a plasmid, cosmid, viral particle, or phage. The appropriate nucleic acid sequence may be inserted into the vector by a variety of procedures. In general, DNA is inserted into an appropriate restriction endonuclease site(s) using techniques known in the art. Vector components generally include, but are not limited to, one or more of a signal sequence, an origin of replication, one or more marker genes, an enhancer element, a promoter, and a transcription termination sequence. Construction of suitable vectors containing one or more of these components employs standard ligation techniques which are known to the skilled artisan.

The PRO may be produced recombinantly not only directly, but also as a fusion polypeptide with a heterologous polypeptide, which may be a signal sequence or other polypeptide having a specific cleavage site at the N-terminus of the mature protein or polypeptide. In general, the signal sequence may be a component of the vector, or it may be a part of the PRO-encoding DNA that is inserted into the vector. The signal sequence may be a prokaryotic signal sequence selected, for example, from the group of the alkaline phosphatase, penicillinase, lpp, or heat-stable enterotoxin II leaders. For yeast secretion the signal sequence may be, e.g., the yeast invertase leader, alpha factor leader (including Saccharomyces and Kluyveromyces are factor leaders, the latter described in U.S. Patent No. 5,010,182), or acid phosphatase leader, the C. albicans glucoamylase leader (EP 362,179 published 4 April 1990), or the signal described in WO 90/13646 published 15 November 1990. In mammalian cell expression, mammalian signal sequences may be used to direct secretion of the protein, such as signal sequences from secreted polypeptides of the same or related species, as well as viral secretory leaders.

Both expression and cloning vectors contain a nucleic acid sequence that enables the vector to replicate in one or more selected host cells. Such sequences are well known for a variety of bacteria, yeast, and viruses. The origin of replication from the plasmid pBR322 is suitable for most Gram-negative bacteria, the 2µ plasmid origin is suitable for yeast, and various viral origins (SV40, polyoma, adenovirus, VSV or BPV) are useful for cloning vectors in mammalian cells.

Expression and cloning vectors will typically contain a selection gene, also termed a selectable marker. Typical selection genes encode proteins that (a) confer resistance to antibiotics or other toxins, e.g., ampicillin, neomycin, methotrexate, or tetracycline, (b) complement auxotrophic deficiencies, or (c) supply critical nutrients not available from complex media, e.g., the gene encoding D-alanine racemase for *Bacilli*.

An example of suitable selectable markers for mammalian cells are those that enable the identification of cells competent to take up the PRO-encoding nucleic acid, such as DHFR or thymidine kinase. An appropriate host cell when wild-type DHFR is employed is the CHO cell line deficient in DHFR

activity, prepared and propagated as described by Urlaub et al., <u>Proc. Natl. Acad. Sci. USA</u>, 77:4216 (1980). A suitable selection gene for use in yeast is the *trp*1 gene present in the yeast plasmid YRp7 [Stinchcomb et al., <u>Nature</u>, 282:39 (1979); Kingsman et al., <u>Gene</u>, 7:141 (1979); Tschemper et al., <u>Gene</u>, 10:157 (1980)]. The *trp*1 gene provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan, for example, ATCC No. 44076 or PEP4-1 [Jones, <u>Genetics</u>, 85:12 (1977)].

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Expression and cloning vectors usually contain a promoter operably linked to the PRO-encoding nucleic acid sequence to direct mRNA synthesis. Promoters recognized by a variety of potential host cells are well known. Promoters suitable for use with prokaryotic hosts include the β-lactamase and lactose promoter systems [Chang et al., Nature, 275:615 (1978); Goeddel et al., Nature, 281:544 (1979)], alkaline phosphatase, a tryptophan (trp) promoter system [Goeddel, Nucleic Acids Res., 8:4057 (1980); EP 36,776], and hybrid promoters such as the tac promoter [deBoer et al., Proc. Natl. Acad. Sci. USA, 80:21-25 (1983)]. Promoters for use in bacterial systems also will contain a Shine-Dalgarno (S.D.) sequence operably linked to the DNA encoding PRO.

Examples of suitable promoting sequences for use with yeast hosts include the promoters for 3-phosphoglycerate kinase [Hitzeman et al., <u>J. Biol. Chem.</u>, 255:2073 (1980)] or other glycolytic enzymes [Hess et al., <u>J. Adv. Enzyme Reg.</u>, 7:149 (1968); Holland, <u>Biochemistry</u>, 17:4900 (1978)], such as enolase, glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucose isomerase, and glucokinase.

Other yeast promoters, which are inducible promoters having the additional advantage of transcription controlled by growth conditions, are the promoter regions for alcohol dehydrogenase 2, isocytochrome C, acid phosphatase, degradative enzymes associated with nitrogen metabolism, metallothionein, glyceraldehyde-3-phosphate dehydrogenase, and enzymes responsible for maltose and galactose utilization. Suitable vectors and promoters for use in yeast expression are further described in EP 73,657.

PRO transcription from vectors in mammalian host cells is controlled, for example, by promoters obtained from the genomes of viruses such as polyoma virus, fowlpox virus (UK 2,211,504 published 5 July 1989), adenovirus (such as Adenovirus 2), bovine papilloma virus, avian sarcoma virus, cytomegalovirus, a retrovirus, hepatitis-B virus and Simian Virus 40 (SV40), from heterologous mammalian promoters, e.g., the actin promoter or an immunoglobulin promoter, and from heat-shock promoters, provided such promoters are compatible with the host cell systems.

Transcription of a DNA encoding the PRO by higher eukaryotes may be increased by inserting an enhancer sequence into the vector. Enhancers are cis-acting elements of DNA, usually about from 10 to 300 bp, that act on a promoter to increase its transcription. Many enhancer sequences are now known from mammalian genes (globin, elastase, albumin, α-fetoprotein, and insulin). Typically, however, one will use an enhancer from a eukaryotic cell virus. Examples include the SV40 enhancer on the late side of the replication origin (bp 100-270), the cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers. The enhancer may be spliced into the vector at a position 5' or 3' to the PRO coding sequence, but is preferably located at a site 5' from the promoter.

Expression vectors used in eukaryotic host cells (yeast, fungi, insect, plant, animal, human, or nucleated cells from other multicellular organisms) will also contain sequences necessary for the termination of transcription and for stabilizing the mRNA. Such sequences are commonly available from the 5' and, occasionally 3', untranslated regions of eukaryotic or viral DNAs or cDNAs. These regions contain nucleotide segments transcribed as polyadenylated fragments in the untranslated portion of the mRNA encoding PRO.

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Still other methods, vectors, and host cells suitable for adaptation to the synthesis of PRO in recombinant vertebrate cell culture are described in Gething et al., Nature, 293:620-625 (1981); Mantei et al., Nature, 281:40-46 (1979); EP 117,060; and EP 117,058.

4. Detecting Gene Amplification/Expression

Gene amplification and/or expression may be measured in a sample directly, for example, by conventional Southern blotting, Northern blotting to quantitate the transcription of mRNA [Thomas, Proc. Natl. Acad. Sci. USA, 77:5201-5205 (1980)], dot blotting (DNA analysis), or in situ hybridization, using an appropriately labeled probe, based on the sequences provided herein. Alternatively, antibodies may be employed that can recognize specific duplexes, including DNA duplexes, RNA duplexes, and DNA-RNA hybrid duplexes or DNA-protein duplexes. The antibodies in turn may be labeled and the assay may be carried out where the duplex is bound to a surface, so that upon the formation of duplex on the surface, the presence of antibody bound to the duplex can be detected.

Gene expression, alternatively, may be measured by immunological methods, such as immunohistochemical staining of cells or tissue sections and assay of cell culture or body fluids, to quantitate directly the expression of gene product. Antibodies useful for immunohistochemical staining and/or assay of sample fluids may be either monoclonal or polyclonal, and may be prepared in any mammal. Conveniently, the antibodies may be prepared against a native sequence PRO polypeptide or against a synthetic peptide based on the DNA sequences provided herein or against exogenous sequence fused to PRO DNA and encoding a specific antibody epitope.

5. Purification of Polypeptide

Forms of PRO may be recovered from culture medium or from host cell lysates. If membrane-bound, it can be released from the membrane using a suitable detergent solution (e.g. Triton-X 100) or by enzymatic cleavage. Cells employed in expression of PRO can be disrupted by various physical or chemical means, such as freeze-thaw cycling, sonication, mechanical disruption, or cell lysing agents.

It may be desired to purify PRO from recombinant cell proteins or polypeptides. The following procedures are exemplary of suitable purification procedures: by fractionation on an ion-exchange column; ethanol precipitation; reverse phase HPLC; chromatography on silica or on a cation-exchange resin such as DEAE; chromatofocusing; SDS-PAGE; ammonium sulfate precipitation; gel filtration using, for example, Sephadex G-75; protein A Sepharose columns to remove contaminants such as IgG; and metal chelating columns to bind epitope-tagged forms of the PRO. Various methods of protein purification may be employed and such methods are known in the art and described for example in Deutscher, Methods in Enzymology, 182 (1990); Scopes, Protein Purification: Principles and Practice, Springer-Verlag, New York (1982). The purification step(s) selected will depend, for example, on the nature of the production process used and the particular PRO produced.

E. <u>Tissue Distribution</u>

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The location of tissues expressing the PRO can be identified by determining mRNA expression in various human tissues. The location of such genes provides information about which tissues are most likely to be affected by the stimulating and inhibiting activities of the PRO polypeptides. The location of a gene in a specific tissue also provides sample tissue for the activity blocking assays discussed below.

As noted before, gene expression in various tissues may be measured by conventional Southern blotting, Northern blotting to quantitate the transcription of mRNA (Thomas, *Proc. Natl. Acad. Sci. USA*, 77:5201-5205 [1980]), dot blotting (DNA analysis), or *in situ* hybridization, using an appropriately labeled probe, based on the sequences provided herein. Alternatively, antibodies may be employed that can recognize specific duplexes, including DNA duplexes, RNA duplexes, and DNA-RNA hybrid duplexes or DNA-protein duplexes.

Gene expression in various tissues, alternatively, may be measured by immunological methods, such as immunohistochemical staining of tissue sections and assay of cell culture or body fluids, to quantitate directly the expression of gene product. Antibodies useful for immunohistochemical staining and/or assay of sample fluids may be either monoclonal or polyclonal, and may be prepared in any mammal. Conveniently, the antibodies may be prepared against a native sequence of a PRO polypeptide or against a synthetic peptide based on the DNA sequences encoding the PRO polypeptide or against an exogenous sequence fused to a DNA encoding a PRO polypeptide and encoding a specific antibody epitope. General techniques for generating antibodies, and special protocols for Northern blotting and *in situ* hybridization are provided below.

F. Antibody Binding Studies

The activity of the PRO polypeptides can be further verified by antibody binding studies, in which the ability of anti-PRO antibodies to inhibit the effect of the PRO polypeptides, respectively, on tissue cells is tested. Exemplary antibodies include polyclonal, monoclonal, humanized, bispecific, and heteroconjugate antibodies, the preparation of which will be described hereinbelow.

Antibody binding studies may be carried out in any known assay method, such as competitive binding assays, direct and indirect sandwich assays, and immunoprecipitation assays. Zola, *Monoclonal Antibodies: A Manual of Techniques*, pp.147-158 (CRC Press, Inc., 1987).

Competitive binding assays rely on the ability of a labeled standard to compete with the test sample analyte for binding with a limited amount of antibody. The amount of target protein in the test sample is inversely proportional to the amount of standard that becomes bound to the antibodies. To facilitate determining the amount of standard that becomes bound, the antibodies preferably are insolubilized before or after the competition, so that the standard and analyte that are bound to the antibodies may conveniently be separated from the standard and analyte which remain unbound.

Sandwich assays involve the use of two antibodies, each capable of binding to a different immunogenic portion, or epitope, of the protein to be detected. In a sandwich assay, the test sample analyte is bound by a first antibody which is immobilized on a solid support, and thereafter a second antibody binds to the analyte, thus forming an insoluble three-part complex. See, e.g., US Pat No. 4,376,110. The second antibody may itself be labeled with a detectable moiety (direct sandwich assays) or may be measured using

an anti-immunoglobulin antibody that is labeled with a detectable moiety (indirect sandwich assay). For example, one type of sandwich assay is an ELISA assay, in which case the detectable moiety is an enzyme.

For immunohistochemistry, the tissue sample may be fresh or frozen or may be embedded in paraffin and fixed with a preservative such as formalin, for example.

G. <u>Cell-Based Assays</u>

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Cell-based assays and animal models for immune related diseases can be used to further understand the relationship between the genes and polypeptides identified herein and the development and pathogenesis of immune related disease.

In a different approach, cells of a cell type known to be involved in a particular immune related disease are transfected with the cDNAs described herein, and the ability of these cDNAs to stimulate or inhibit immune function is analyzed. Suitable cells can be transfected with the desired gene, and monitored for immune function activity. Such transfected cell lines can then be used to test the ability of poly- or monoclonal antibodies or antibody compositions to inhibit or stimulate immune function, for example to modulate T-cell proliferation or inflammatory cell infiltration. Cells transfected with the coding sequences of the genes identified herein can further be used to identify drug candidates for the treatment of immune related diseases.

In addition, primary cultures derived from transgenic animals (as described below) can be used in the cell-based assays herein, although stable cell lines are preferred. Techniques to derive continuous cell lines from transgenic animals are well known in the art (see, e.g., Small et al., Mol. Cell. Biol. 5: 642-648 [1985]).

One suitable cell based assay is the mixed lymphocyte reaction (MLR). Current Protocols in Immunology, unit 3.12; edited by J E Coligan, A M Kruisbeek, D H Marglies, E M Shevach, W Strober, National Institutes of Health, Published by John Wiley & Sons, Inc. In this assay, the ability of a test compound to stimulate or inhibit the proliferation of activated T cells is assayed. A suspension of responder T cells is cultured with allogeneic stimulator cells and the proliferation of T cells is measured by uptake of tritiated thymidine. This assay is a general measure of T cell reactivity. Since the majority of T cells respond to and produce IL-2 upon activation, differences in responsiveness in this assay in part reflect differences in IL-2 production by the responding cells. The MLR results can be verified by a standard lymphokine (IL-2) detection assay. Current Protocols in Immunology, above, 3.15, 6.3.

A proliferative T cell response in an MLR assay may be due to direct mitogenic properties of an assayed molecule or to external antigen induced activation. Additional verification of the T cell stimulatory activity of the PRO polypeptides can be obtained by a costimulation assay. T cell activation requires an antigen specific signal mediated through the T-cell receptor (TCR) and a costimulatory signal mediated through a second ligand binding interaction, for example, the B7 (CD80, CD86)/CD28 binding interaction. CD28 crosslinking increases lymphokine secretion by activated T cells. T cell activation has both negative and positive controls through the binding of ligands which have a negative or positive effect. CD28 and CTLA-4 are related glycoproteins in the Ig superfamily which bind to B7. CD28 binding to B7 has a positive costimulation effect of T cell activation; conversely, CTLA-4 binding to B7 has a T cell deactivating effect. Chambers, C. A. and Allison, J. P., Curr. Opin. Immunol. (1997) 2:396. Schwartz, R. H., Cell (1992) 71:1065; Linsey, P. S. and Ledbetter, J. A., Annu. Rev. Immunol. (1993) 11:191; June, C. H.

et al, Immunol. Today (1994) 15:321; Jenkins, M. K., Immunity (1994) 1:405. In a costimulation assay, the PRO polypeptides are assayed for T cell costimulatory or inhibitory activity.

Direct use of a stimulating compound as in the invention has been validated in experiments with 4-1BB glycoprotein, a member of the tumor necrosis factor receptor family, which binds to a ligand (4-1BBL) expressed on primed T cells and signals T cell activation and growth. Alderson, M. E. et al., J. Immunol. (1994) 24:2219.

The use of an agonist stimulating compound has also been validated experimentally. Activation of 4-1BB by treatment with an agonist anti-4-1BB antibody enhances eradication of tumors. Hellstrom, I. and Hellstrom, K. E., *Crit. Rev. Immunol.* (1998) 18:1. Immunoadjuvant therapy for treatment of tumors, described in more detail below, is another example of the use of the stimulating compounds of the invention.

Alternatively, an immune stimulating or enhancing effect can also be achieved by administration of a PRO which has vascular permeability enhancing properties. Enhanced vascular permeability would be beneficial to disorders which can be attenuated by local infiltration of immune cells (e.g., monocytes, eosinophils, PMNs) and inflammation.

On the other hand, PRO polypeptides, as well as other compounds of the invention, which are direct inhibitors of T cell proliferation/activation, lymphokine secretion, and/or vascular permeability can be directly used to suppress the immune response. These compounds are useful to reduce the degree of the immune response and to treat immune related diseases characterized by a hyperactive, superoptimal, or autoimmune response. This use of the compounds of the invention has been validated by the experiments described above in which CTLA-4 binding to receptor B7 deactivates T cells. The direct inhibitory compounds of the invention function in an analogous manner. The use of compound which suppress vascular permeability would be expected to reduce inflammation. Such uses would be beneficial in treating conditions associated with excessive inflammation.

Alternatively, compounds, e.g., antibodies, which bind to stimulating PRO polypeptides and block the stimulating effect of these molecules produce a net inhibitory effect and can be used to suppress the T cell mediated immune response by inhibiting T cell proliferation/activation and/or lymphokine secretion. Blocking the stimulating effect of the polypeptides suppresses the immune response of the mammal. This use has been validated in experiments using an anti-IL2 antibody. In these experiments, the antibody binds to IL2 and blocks binding of IL2 to its receptor thereby achieving a T cell inhibitory effect.

H. Animal Models

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The results of the cell based in vitro assays can be further verified using in vivo animal models and assays for T-cell function. A variety of well known animal models can be used to further understand the role of the genes identified herein in the development and pathogenesis of immune related disease, and to test the efficacy of candidate therapeutic agents, including antibodies, and other antagonists of the native polypeptides, including small molecule antagonists. The in vivo nature of such models makes them predictive of responses in human patients. Animal models of immune related diseases include both non-recombinant and recombinant (transgenic) animals. Non-recombinant animal models include, for example, rodent, e.g., murine models. Such models can be generated by introducing cells into syngeneic mice using

standard techniques, e.g., subcutaneous injection, tail vein injection, spleen implantation, intraperitoneal implantation, implantation under the renal capsule, etc.

Graft-versus-host disease occurs when immunocompetent cells are transplanted into immunosuppressed or tolerant patients. The donor cells recognize and respond to host antigens. The response can vary from life threatening severe inflammation to mild cases of diarrhea and weight loss. Graft-versus-host disease models provide a means of assessing T cell reactivity against MHC antigens and minor transplant antigens. A suitable procedure is described in detail in Current Protocols in Immunology, above, unit 4.3.

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An animal model for skin allograft rejection is a means of testing the ability of T cells to mediate in vivo tissue destruction and a measure of their role in transplant rejection. The most common and accepted models use murine tail-skin grafts. Repeated experiments have shown that skin allograft rejection is mediated by T cells, helper T cells and killer-effector T cells, and not antibodies. Auchincloss, H. Jr. and Sachs, D. H., Fundamental Immunology, 2nd ed., W. E. Paul ed., Raven Press, NY, 1989, 889-992. A suitable procedure is described in detail in Current Protocols in Immunology, above, unit 4.4. Other transplant rejection models which can be used to test the compounds of the invention are the allogeneic heart transplant models described by Tanabe, M. et al, Transplantation (1994) 58:23 and Tinubu, S. A. et al, J. Immunol. (1994) 4330-4338.

Animal models for delayed type hypersensitivity provides an assay of cell mediated immune function as well. Delayed type hypersensitivity reactions are a T cell mediated in vivo immune response characterized by inflammation which does not reach a peak until after a period of time has elapsed after challenge with an antigen. These reactions also occur in tissue specific autoimmune diseases such as multiple sclerosis (MS) and experimental autoimmune encephalomyelitis (EAE, a model for MS). A suitable procedure is described in detail in Current Protocols in Immunology, above, unit 4.5.

EAE is a T cell mediated autoimmune disease characterized by T cell and mononuclear cell inflammation and subsequent demyelination of axons in the central nervous system. EAE is generally considered to be a relevant animal model for MS in humans. Bolton, C., *Multiple Sclerosis* (1995) 1:143. Both acute and relapsing-remitting models have been developed. The compounds of the invention can be tested for T cell stimulatory or inhibitory activity against immune mediated demyelinating disease using the protocol described in *Current Protocols in Immunology*, above, units 15.1 and 15.2. See also the models for myelin disease in which oligodendrocytes or Schwann cells are grafted into the central nervous system as described in Duncan, I. D. *et al*, *Molec. Med. Today* (1997) 554-561.

Contact hypersensitivity is a simple delayed type hypersensitivity in vivo assay of cell mediated immune function. In this procedure, cutaneous exposure to exogenous haptens which gives rise to a delayed type hypersensitivity reaction which is measured and quantitated. Contact sensitivity involves an initial sensitizing phase followed by an elicitation phase. The elicitation phase occurs when the T lymphocytes encounter an antigen to which they have had previous contact. Swelling and inflammation occur, making this an excellent model of human allergic contact dermatitis. A suitable procedure is described in detail in Current Protocols in Immunology, Eds. J. E. Cologan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach and W. Strober, John Wiley & Sons, Inc., 1994, unit 4.2. See also Grabbe, S. and Schwarz, T, Immun. Today 19 (1): 37-44 (1998).

An animal model for arthritis is collagen-induced arthritis. This model shares clinical, histological and immunological characteristics of human autoimmune rheumatoid arthritis and is an acceptable model for human autoimmune arthritis. Mouse and rat models are characterized by synovitis, erosion of cartilage and subchondral bone. The compounds of the invention can be tested for activity against autoimmune arthritis using the protocols described in *Current Protocols in Immunology*, above, units 15.5. See also the model using a monoclonal antibody to CD18 and VLA-4 integrins described in Issekutz, A.C. et al., Immunology (1996) 88:569.

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A model of asthma has been described in which antigen-induced airway hyper-reactivity, pulmonary eosinophilia and inflammation are induced by sensitizing an animal with ovalbumin and then challenging the animal with the same protein delivered by aerosol. Several animal models (guinea pig, rat, non-human primate) show symptoms similar to atopic asthma in humans upon challenge with aerosol antigens. Murine models have many of the features of human asthma. Suitable procedures to test the compounds of the invention for activity and effectiveness in the treatment of asthma are described by Wolyniec, W. W. et al, Am. J. Respir. Cell Mol. Biol. (1998) 18:777 and the references cited therein.

Additionally, the compounds of the invention can be tested on animal models for 'psoriasis like diseases. Evidence suggests a T cell pathogenesis for psoriasis. The compounds of the invention can be tested in the scid/scid mouse model described by Schon, M. P. et al, Nat. Med. (1997) 3:183, in which the mice demonstrate histopathologic skin lesions resembling psoriasis. Another suitable model is the human skin/scid mouse chimera prepared as described by Nickoloff, B. J. et al, Am. J. Path. (1995) 146:580.

Recombinant (transgenic) animal models can be engineered by introducing the coding portion of the genes identified herein into the genome of animals of interest, using standard techniques for producing transgenic animals. Animals that can serve as a target for transgenic manipulation include, without limitation, mice, rats, rabbits, guinea pigs, sheep, goats, pigs, and non-human primates, e.g., baboons, chimpanzees and monkeys. Techniques known in the art to introduce a transgene into such animals include pronucleic microinjection (Hoppe and Wanger, U.S. Patent No. 4,873,191); retrovirus-mediated gene transfer into germ lines (e.g., Van der Putten et al., Proc. Natl. Acad. Sci. USA 82, 6148-615 [1985]); gene targeting in embryonic stem cells (Thompson et al., Cell 56, 313-321 [1989]); electroporation of embryos (Lo, Mol. Cel. Biol. 3, 1803-1814 [1983]); sperm-mediated gene transfer (Lavitrano et al., Cell 57, 717-73 [1989]). For review, see, for example, U.S. Patent No. 4,736,866.

For the purpose of the present invention, transgenic animals include those that carry the transgene only in part of their cells ("mosaic animals"). The transgene can be integrated either as a single transgene, or in concatamers, e.g., head-to-head or head-to-tail tandems. Selective introduction of a transgene into a particular cell type is also possible by following, for example, the technique of Lasko et al., Proc. Natl. Acad. Sci. USA 89, 6232-636 (1992).

The expression of the transgene in transgenic animals can be monitored by standard techniques. For example, Southern blot analysis or PCR amplification can be used to verify the integration of the transgene. The level of mRNA expression can then be analyzed using techniques such as *in situ* hybridization, Northern blot analysis, PCR, or immunocytochemistry.

The animals may be further examined for signs of immune disease pathology, for example by histological examination to determine infiltration of immune cells into specific tissues. Blocking

experiments can also be performed in which the transgenic animals are treated with the compounds of the invention to determine the extent of the T cell proliferation stimulation or inhibition of the compounds. In these experiments, blocking antibodies which bind to the PRO polypeptide, prepared as described above, are administered to the animal and the effect on immune function is determined.

Alternatively, "knock out" animals can be constructed which have a defective or altered gene encoding a polypeptide identified herein, as a result of homologous recombination between the endogenous . gene encoding the polypeptide and altered genomic DNA encoding the same polypeptide introduced into an embryonic cell of the animal. For example, cDNA encoding a particular polypeptide can be used to clone genomic DNA encoding that polypeptide in accordance with established techniques. A portion of the genomic DNA encoding a particular polypeptide can be deleted or replaced with another gene, such as a gene encoding a selectable marker which can be used to monitor integration. Typically, several kilobases of unaltered flanking DNA (both at the 5' and 3' ends) are included in the vector [see e.g., Thomas and Capecchi, Cell, 51:503 (1987) for a description of homologous recombination vectors]. The vector is introduced into an embryonic stem cell line (e.g., by electroporation) and cells in which the introduced DNA has homologously recombined with the endogenous DNA are selected [see e.g., Li et al., Cell, 69:915 (1992)]. The selected cells are then injected into a blastocyst of an animal (e.g., a mouse or rat) to form aggregation chimeras [see e.g., Bradley, in Teratocarcinomas and Embryonic Stem Cells: A Practical Approach, E. J. Robertson, ed. (IRL, Oxford, 1987), pp. 113-152]. A chimeric embryo can then be implanted into a suitable pseudopregnant female foster animal and the embryo brought to term to create a "knock out" animal. Progeny harboring the homologously recombined DNA in their germ cells can be identified by standard techniques and used to breed animals in which all cells of the animal contain the homologously recombined DNA. Knockout animals can be characterized for instance, for their ability to defend against certain pathological conditions and for their development of pathological conditions due to absence of the polypeptide.

I. ImmunoAdjuvant Therapy

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In one embodiment, the immunostimulating compounds of the invention can be used in immunoadjuvant therapy for the treatment of tumors (cancer). It is now well established that T cells recognize human tumor specific antigens. One group of tumor antigens, encoded by the MAGE, BAGE and GAGE families of genes, are silent in all adult normal tissues, but are expressed in significant amounts in tumors, such as melanomas, lung tumors, head and neck tumors, and bladder carcinomas. DeSmet, C. et al., (1996) Proc. Natl. Acad. Sci. USA, 93:7149. It has been shown that costimulation of T cells induces tumor regression and an antitumor response both in vitro and in vivo. Melero, I. et al., Nature Medicine (1997) 3:682; Kwon, E. D. et al., Proc. Natl. Acad. Sci. USA (1997) 94: 8099; Lynch, D. H. et al, Nature Medicine (1997) 3:625; Finn, O. J. and Lotze, M. T., J. Immunol. (1998) 21:114. The stimulatory compounds of the invention can be administered as adjuvants, alone or together with a growth regulating agent, cytotoxic agent or chemotherapeutic agent, to stimulate T cell proliferation/activation and an antitumor response to tumor antigens. The growth regulating, cytotoxic, or chemotherapeutic agent may be administered in conventional amounts using known administration regimes. Immunostimulating activity by the compounds of the invention allows reduced amounts of the growth regulating, cytotoxic, or chemotherapeutic agents thereby potentially lowering the toxicity to the patient.

J. Screening Assays for Drug Candidates

Screening assays for drug candidates are designed to identify compounds that bind to or complex with the polypeptides encoded by the genes identified herein or a biologically active fragment thereof, or otherwise interfere with the interaction of the encoded polypeptides with other cellular proteins. Such screening assays will include assays amenable to high-throughput screening of chemical libraries, making them particularly suitable for identifying small molecule drug candidates. Small molecules contemplated include synthetic organic or inorganic compounds, including peptides, preferably soluble peptides, (poly)peptide-immunoglobulin fusions, and, in particular, antibodies including, without limitation, poly- and monoclonal antibodies and antibody fragments, single-chain antibodies, anti-idiotypic antibodies, and chimeric or humanized versions of such antibodies or fragments, as well as human antibodies and antibody fragments. The assays can be performed in a variety of formats, including protein-protein binding assays, biochemical screening assays, immunoassays and cell based assays, which are well characterized in the art. All assays are common in that they call for contacting the drug candidate with a polypeptide encoded by a nucleic acid identified herein under conditions and for a time sufficient to allow these two components to interact.

In binding assays, the interaction is binding and the complex formed can be isolated or detected in the reaction mixture. In a particular embodiment, the polypeptide encoded by the gene identified herein or the drug candidate is immobilized on a solid phase, e.g., on a microtiter plate, by covalent or non-covalent attachments. Non-covalent attachment generally is accomplished by coating the solid surface with a solution of the polypeptide and drying. Alternatively, an immobilized antibody, e.g., a monoclonal antibody, specific for the polypeptide to be immobilized can be used to anchor it to a solid surface. The assay is performed by adding the non-immobilized component, which may be labeled by a detectable label, to the immobilized component, e.g., the coated surface containing the anchored component. When the reaction is complete, the non-reacted components are removed, e.g., by washing, and complexes anchored on the solid surface are detected. When the originally non-immobilized component carries a detectable label, the detection of label immobilized on the surface indicates that complexing occurred. Where the originally non-immobilized component does not carry a label, complexing can be detected, for example, by using a labelled antibody specifically binding the immobilized complex.

If the candidate compound interacts with but does not bind to a particular protein encoded by a gene identified herein, its interaction with that protein can be assayed by methods well known for detecting protein-protein interactions. Such assays include traditional approaches, such as, cross-linking, co-immunoprecipitation, and co-purification through gradients or chromatographic columns. In addition, protein-protein interactions can be monitored by using a yeast-based genetic system described by Fields and co-workers [Fields and Song, Nature (London) 340, 245-246 (1989); Chien et al., Proc. Natl. Acad. Sci. USA 88, 9578-9582 (1991)] as disclosed by Chevray and Nathans, Proc. Natl. Acad. Sci. USA 89, 5789-5793 (1991). Many transcriptional activators, such as yeast GALA, consist of two physically discrete modular domains, one acting as the DNA-binding domain, while the other one functioning as the transcription activation domain. The yeast expression system described in the foregoing publications (generally referred to as the "two-hybrid system") takes advantage of this property, and employs two hybrid proteins, one in which the target protein is fused to the DNA-binding domain of GALA, and another, in which candidate

activating proteins are fused to the activation domain. The expression of a GAL1-lacZ reporter gene under control of a GAL4-activated promoter depends on reconstitution of GAL4 activity via protein-protein interaction. Colonies containing interacting polypeptides are detected with a chromogenic substrate for β-galactosidase. A complete kit (MATCHMAKERTM) for identifying protein-protein interactions between two specific proteins using the two-hybrid technique is commercially available from Clontech. This system can also be extended to map protein domains involved in specific protein interactions as well as to pinpoint amino acid residues that are crucial for these interactions.

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In order to find compounds that interfere with the interaction of a gene identified herein and other intra- or extracellular components can be tested, a reaction mixture is usually prepared containing the product of the gene and the intra- or extracellular component under conditions and for a time allowing for the interaction and binding of the two products. To test the ability of a test compound to inhibit binding, the reaction is run in the absence and in the presence of the test compound. In addition, a placebo may be added to a third reaction mixture, to serve as positive control. The binding (complex formation) between the test compound and the intra- or extracellular component present in the mixture is monitored as described above. The formation of a complex in the control reaction(s) but not in the reaction mixture containing the test compound indicates that the test compound interferes with the interaction of the test compound and its reaction partner.

K. Compositions and Methods for the Treatment of Immune Related Diseases

The compositions useful in the treatment of immune related diseases include, without limitation, proteins, antibodies, small organic molecules, peptides, phosphopeptides, antisense and ribozyme molecules, triple helix molecules, etc. that inhibit or stimulate immune function, for example, T cell proliferation/activation, lymphokine release, or immune cell infiltration.

For example, antisense RNA and RNA molecules act to directly block the translation of mRNA by hybridizing to targeted mRNA and preventing protein translation. When antisense DNA is used, oligodeoxyribonucleotides derived from the translation initiation site, e.g., between about -10 and +10 positions of the target gene nucleotide sequence, are preferred.

Ribozymes are enzymatic RNA molecules capable of catalyzing the specific cleavage of RNA. Ribozymes act by sequence-specific hybridization to the complementary target RNA, followed by endonucleolytic cleavage. Specific ribozyme cleavage sites within a potential RNA target can be identified by known techniques. For further details see, e.g., Rossi, Current Biology 4, 469-471 (1994), and PCT publication No. WO 97/33551 (published September 18, 1997).

Nucleic acid molecules in triple helix formation used to inhibit transcription should be single-stranded and composed of deoxynucleotides. The base composition of these oligonucleotides is designed such that it promotes triple helix formation via Hoogsteen base pairing rules, which generally require sizeable stretches of purines or pyrimidines on one strand of a duplex. For further details see, e.g., PCT publication No. WO 97/33551, supra.

These molecules can be identified by any or any combination of the screening assays discussed above and/or by any other screening techniques well known for those skilled in the art.

L. Anti-PRO Antibodies

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The present invention further provides anti-PRO antibodies. Exemplary antibodies include polyclonal, monoclonal, humanized, bispecific, and heteroconjugate antibodies.

1. Polyclonal Antibodies

The anti-PRO antibodies may comprise polyclonal antibodies. Methods of preparing polyclonal antibodies are known to the skilled artisan. Polyclonal antibodies can be raised in a mammal, for example, by one or more injections of an immunizing agent and, if desired, an adjuvant. Typically, the immunizing agent and/or adjuvant will be injected in the mammal by multiple subcutaneous or intraperitoneal injections. The immunizing agent may include the PRO polypeptide or a fusion protein thereof. It may be useful to conjugate the immunizing agent to a protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. Examples of adjuvants which may be employed include Freund's complete adjuvant and MPL-TDM adjuvant (monophosphoryl Lipid A, synthetic trehalose dicorynomycolate). The immunization protocol may be selected by one skilled in the art without undue experimentation.

2. Monoclonal Antibodies

The anti-PRO antibodies may, alternatively, be monoclonal antibodies. Monoclonal antibodies may be prepared using hybridoma methods, such as those described by Kohler and Milstein, Nature, 256:495 (1975). In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes may be immunized in vitro.

The immunizing agent will typically include the PRO polypeptide or a fusion protein thereof. Generally, either peripheral blood lymphocytes ("PBLs") are used if cells of human origin are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell [Goding, Monoclonal Antibodies: Principles and Practice, Academic Press, (1986) pp. 59-103]. Immortalized cell lines are usually transformed mammalian cells, particularly myeloma cells of rodent, bovine and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells may be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

Preferred immortalized cell lines are those that fuse efficiently, support stable high level expression of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. More preferred immortalized cell lines are murine myeloma lines, which can be obtained, for instance, from the Salk Institute Cell Distribution Center, San Diego, California and the American Type Culture Collection, Manassas, Virginia. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies [Kozbor, J. Immunol., 133:3001 (1984); Brodeur et al.,

Monoclonal Antibody Production Techniques and Applications, Marcel Dekker, Inc., New York, (1987) pp. 51-63].

The culture medium in which the hybridoma cells are cultured can then be assayed for the presence of monoclonal antibodies directed against PRO. Preferably, the binding specificity of monoclonal antibodies produced by the hybridoma cells is determined by immunoprecipitation or by an *in vitro* binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent assay (ELISA). Such techniques and assays are known in the art. The binding affinity of the monoclonal antibody can, for example, be determined by the Scatchard analysis of Munson and Pollard, Anal. Biochem., 107:220 (1980).

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After the desired hybridoma cells are identified, the clones may be subcloned by limiting dilution procedures and grown by standard methods [Goding, <u>supra</u>]. Suitable culture media for this purpose include, for example, Dulbecco's Modified Eagle's Medium and RPMI-1640 medium. Alternatively, the hybridoma cells may be grown *in vivo* as ascites in a mammal.

The monoclonal antibodies secreted by the subclones may be isolated or purified from the culture medium or ascites fluid by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography.

The monoclonal antibodies may also be made by recombinant DNA methods, such as those described in U.S. Patent No. 4,816,567. DNA encoding the monoclonal antibodies of the invention can be readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of murine antibodies). The hybridoma cells of the invention serve as a preferred source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as simian COS cells, Chinese hamster ovary (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. The DNA also may be modified, for example, by substituting the coding sequence for human heavy and light chain constant domains in place of the homologous murine sequences [U.S. Patent No. 4,816,567; Morrison et al., supra] or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence for a non-immunoglobulin polypeptide. Such a non-immunoglobulin polypeptide can be substituted for the constant domains of an antibody of the invention, or can be substituted for the variable domains of one antigencombining site of an antibody of the invention to create a chimeric bivalent antibody.

The antibodies may be monovalent antibodies. Methods for preparing monovalent antibodies are well known in the art. For example, one method involves recombinant expression of immunoglobulin light chain and modified heavy chain. The heavy chain is truncated generally at any point in the Fc region so as to prevent heavy chain crosslinking. Alternatively, the relevant cysteine residues are substituted with another amino acid residue or are deleted so as to prevent crosslinking.

In vitro methods are also suitable for preparing monovalent antibodies. Digestion of antibodies to produce fragments thereof, particularly, Fab fragments, can be accomplished using routine techniques known in the art.

3. Human and Humanized Antibodies

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The anti-PRO antibodies of the invention may further comprise humanized antibodies or human antibodies. Humanized forms of non-human (e.g., murine) antibodies are chimeric immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')2 or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. Humanized antibodies include human immunoglobulins (recipient antibody) in which residues from a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a nonhuman species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin [Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-329 (1988); and Presta, Curr. Op. Struct. Biol., 2:593-596 (1992)].

Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source which is non-human. These non-human amino acid residues are often referred to as "import" residues, which are typically taken from an "import" variable domain. Humanization can be essentially performed following the method of Winter and co-workers [Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); Verhoeyen et al., Science, 239:1534-1536 (1988)], by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such "humanized" antibodies are chimeric antibodies (U.S. Patent No. 4,816,567), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues and possibly some FR residues are substituted by residues from analogous sites in rodent antibodies.

Human antibodies can also be produced using various techniques known in the art, including phage display libraries [Hoogenboom and Winter, J. Mol. Biol., 227:381 (1991); Marks et al., J. Mol. Biol., 222:581 (1991)]. The techniques of Cole et al. and Boerner et al. are also available for the preparation of human monoclonal antibodies (Cole et al., Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, p. 77 (1985) and Boerner et al., J. Immunol., 147(1):86-95 (1991)]. Similarly, human antibodies can be made by introducing of human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in all respects, including gene rearrangement, assembly, and antibody repertoire. This approach is described, for example, in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in the following scientific publications: Marks et al., Bio/Technology 10, 779-783 (1992); Lonberg et al., Nature 368 856-859 (1994);

Morrison, Nature 368, 812-13 (1994); Fishwild et al., Nature Biotechnology 14, 845-51 (1996); Neuberger, Nature Biotechnology 14, 826 (1996); Lonberg and Huszar, Intern. Rev. Immunol. 13 65-93 (1995).

The antibodies may also be affinity matured using known selection and/or mutagenesis methods as described above. Preferred affinity matured antibodies have an affinity which is five times, more preferably 10 times, even more preferably 20 or 30 times greater than the starting antibody (generally murine, humanized or human) from which the matured antibody is prepared.

4. Bispecific Antibodies

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Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. In the present case, one of the binding specificities is for the PRO, the other one is for any other antigen, and preferably for a cell-surface protein or receptor or receptor subunit.

Methods for making bispecific antibodies are known in the art. Traditionally, the recombinant production of bispecific antibodies is based on the co-expression of two immunoglobulin heavy-chain/light-chain pairs, where the two heavy chains have different specificities [Milstein and Cuello, Nature, 305:537-539 (1983)]. Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of ten different antibody molecules, of which only one has the correct bispecific structure. The purification of the correct molecule is usually accomplished by affinity chromatography steps. Similar procedures are disclosed in WO 93/08829, published 13 May 1993, and in Traunecker et al., EMBO J., 10:3655-3659 (1991).

Antibody variable domains with the desired binding specificities (antibody-antigen combining sites) can be fused to immunoglobulin constant domain sequences. The fusion preferably is with an immunoglobulin heavy-chain constant domain, comprising at least part of the hinge, CH2, and CH3 regions. It is preferred to have the first heavy-chain constant region (CH1) containing the site necessary for light-chain binding present in at least one of the fusions. DNAs encoding the immunoglobulin heavy-chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and are co-transfected into a suitable host organism. For further details of generating bispecific antibodies see, for example, Suresh et al., Methods in Enzymology, 121:210 (1986).

According to another approach described in WO 96/27011, the interface between a pair of antibody molecules can be engineered to maximize the percentage of heterodimers which are recovered from recombinant cell culture. The preferred interface comprises at least a part of the CH3 region of an antibody constant domain. In this method, one or more small amino acid side chains from the interface of the first antibody molecule are replaced with larger side chains (e.g. tyrosine or tryptophan). Compensatory "cavities" of identical or similar size to the large side chain(s) are created on the interface of the second antibody molecule by replacing large amino acid side chains with smaller ones (e.g. alanine or threonine). This provides a mechanism for increasing the yield of the heterodimer over other unwanted end-products such as homodimers.

Bispecific antibodies can be prepared as full length antibodies or antibody fragments (e.g. F(ab')₂ bispecific antibodies). Techniques for generating bispecific antibodies from antibody fragments have been described in the literature. For example, bispecific antibodies can be prepared can be prepared using

chemical linkage. Brennan et al., Science 229:81 (1985) describe a procedure wherein intact antibodies are proteolytically cleaved to generate F(ab')₂ fragments. These fragments are reduced in the presence of the dithiol complexing agent sodium arsenite to stabilize vicinal dithiols and prevent intermolecular disulfide formation. The Fab' fragments generated are then converted to thionitrobenzoate (TNB) derivatives. One of the Fab'-TNB derivatives is then reconverted to the Fab'-thiol by reduction with mercaptoethylamine and is mixed with an equimolar amount of the other Fab'-TNB derivative to form the bispecific antibody. The bispecific antibodies produced can be used as agents for the selective immobilization of enzymes.

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Fab' fragments may be directly recovered from *E. coli* and chemically coupled to form bispecific antibodies. Shalaby *et al.*, J. Exp. Med. 175:217-225 (1992) describe the production of a fully humanized bispecific antibody F(ab')₂ molecule. Each Fab' fragment was separately secreted from *E. coli* and subjected to directed chemical coupling *in vitro* to form the bispecific antibody. The bispecific antibody thus formed was able to bind to cells overexpressing the ErbB2 receptor and normal human T cells, as well as trigger the lytic activity of human cytotoxic lymphocytes against human breast tumor targets.

Various technique for making and isolating bispecific antibody fragments directly from recombinant cell culture have also been described. For example, bispecific antibodies have been produced using leucine zippers. Kostelny et al., J. Immunol. 148(5):1547-1553 (1992). The leucine zipper peptides from the Fos and Jun proteins were linked to the Fab' portions of two different antibodies by gene fusion. The antibody homodimers were reduced at the hinge region to form monomers and then re-oxidized to form the antibody heterodimers. This method can also be utilized for the production of antibody homodimers. The "diabody" technology described by Hollinger et al., Proc. Natl. Acad. Sci. USA 90:6444-6448 (1993) has provided an alternative mechanism for making bispecific antibody fragments. The fragments comprise a heavy-chain variable domain (V_H) connected to a light-chain variable domain (V_L) by a linker which is too short to allow pairing between the two domains on the same chain. Accordingly, the V_H and V_L domains of one fragment are forced to pair with the complementary V_L and V_H domains of another fragment, thereby forming two antigen-binding sites. Another strategy for making bispecific antibody fragments by the use of single-chain Fv (sFv) dimers has also been reported. See, Gruber et al., J. Immunol. 152:5368 (1994). Antibodies with more than two valencies are contemplated. For example, trispecific antibodies can be prepared. Tutt et al., J. Immunol. 147:60 (1991).

Exemplary bispecific antibodies may bind to two different epitopes on a given PRO polypeptide herein. Alternatively, an anti-PRO polypeptide arm may be combined with an arm which binds to a triggering molecule on a leukocyte such as a T-cell receptor molecule (e.g. CD2, CD3, CD28, or B7), or Fc receptors for IgG (FcγR), such as FcγRI (CD64), FcγRII (CD32) and FcγRIII (CD16) so as to focus cellular defense mechanisms to the cell expressing the particular PRO polypeptide. Bispecific antibodies may also be used to localize cytotoxic agents to cells which express a particular PRO polypeptide. These antibodies possess a PRO-binding arm and an arm which binds a cytotoxic agent or a radionuclide chelator, such as EOTUBE, DPTA, DOTA, or TETA. Another bispecific antibody of interest binds the PRO polypeptide and further binds tissue factor (TF).

5. <u>Heteroconjugate Antibodies</u>

Heteroconjugate antibodies are also within the scope of the present invention. Heteroconjugate antibodies are composed of two covalently joined antibodies. Such antibodies have, for example, been

proposed to target immune system cells to unwanted cells [U.S. Patent No. 4,676,980], and for treatment of HIV infection [WO 91/00360; WO 92/200373; EP 03089]. It is contemplated that the antibodies may be prepared *in vitro* using known methods in synthetic protein chemistry, including those involving crosslinking agents. For example, immunotoxins may be constructed using a disulfide exchange reaction or by forming a thioether bond. Examples of suitable reagents for this purpose include iminothiolate and methyl-4-mercaptobutyrimidate and those disclosed, for example, in U.S. Patent No. 4,676,980.

6. <u>Bffector Function Engineering</u>

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It may be desirable to modify the antibody of the invention with respect to effector function, so as to enhance, e.g., the effectiveness of the antibody in treating cancer. For example, cysteine residue(s) may be introduced into the Fc region, thereby allowing interchain disulfide bond formation in this region. The homodimeric antibody thus generated may have improved internalization capability and/or increased complement-mediated cell killing and antibody-dependent cellular cytotoxicity (ADCC). See Caron et al., J. Exp Med., 176: 1191-1195 (1992) and Shopes, J. Immunol., 148: 2918-2922 (1992). Homodimeric antibodies with enhanced anti-tumor activity may also be prepared using heterobifunctional cross-linkers as described in Wolff et al. Cancer Research, 53: 2560-2565 (1993). Alternatively, an antibody can be engineered that has dual Fc regions and may thereby have enhanced complement lysis and ADCC capabilities. See Stevenson et al., Anti-Cancer Drug Design, 3: 219-230 (1989).

7. <u>Immunoconjugates</u>

The invention also pertains to immunoconjugates comprising an antibody conjugated to a cytotoxic agent such as a chemotherapeutic agent, toxin (e.g., an enzymatically active toxin of bacterial, fungal, plant, or animal origin, or fragments thereof), or a radioactive isotope (i.e., a radioconjugate).

Chemotherapeutic agents useful in the generation of such immunoconjugates have been described above. Enzymatically active toxins and fragments thereof that can be used include diphtheria A chain, nonbinding active fragments of diphtheria toxin, exotoxin A chain (from *Pseudomonas aeruginosa*), ricin A chain, abrin A chain, modeccin A chain, alpha-sarcin, *Aleurites fordii* proteins, dianthin proteins, *Phytolaca americana* proteins (PAPI, PAPII, and PAP-S), momordica charantia inhibitor, curcin, crotin, sapaonaria officinalis inhibitor, gelonin, mitogellin, restrictocin, phenomycin, enomycin, and the tricothecenes. A variety of radionuclides are available for the production of radioconjugated antibodies. Examples include ²¹²Bi, ¹³¹I, ¹³¹In, ⁹⁰Y, and ¹⁸⁶Re.

Conjugates of the antibody and cytotoxic agent are made using a variety of bifunctional protein-coupling agents such as N-succinimidyl-3-(2-pyridyldithiol) propionate (SPDP), iminothiolane (IT), bifunctional derivatives of imidoesters (such as dimethyl adipimidate HCL), active esters (such as disuccinimidyl suberate), aldehydes (such as glutareldehyde), bis-azido compounds (such as bis (p-azidobenzoyl) hexanediamine), bis-diazonium derivatives (such as bis-(p-diazoniumbenzoyl)-ethylenediamine), diisocyanates (such as tolyene 2,6-diisocyanate), and bis-active fluorine compounds (such as 1,5-difluoro-2,4-dinitrobenzene). For example, a ricin immunotoxin can be prepared as described in Vitetta et al., Science, 238: 1098 (1987). Carbon-14-labeled 1-isothiocyanatobenzyl-3-methyldiethylene triaminepentaacetic acid (MX-DTPA) is an exemplary chelating agent for conjugation of radionucleotide to the antibody. See WO94/11026.

In another embodiment, the antibody may be conjugated to a "receptor" (such streptavidin) for utilization in tumor pretargeting wherein the antibody-receptor conjugate is administered to the patient, followed by removal of unbound conjugate from the circulation using a clearing agent and then administration of a "ligand" (e.g., avidin) that is conjugated to a cytotoxic agent (e.g., a radionucleotide).

8. <u>Immunoliposomes</u>

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The antibodies disclosed herein may also be formulated as immunoliposomes. Liposomes containing the antibody are prepared by methods known in the art, such as described in Epstein *et al.*, <u>Proc. Natl. Acad. Sci. USA</u>, <u>82</u>: 3688 (1985); Hwang *et al.*, <u>Proc. Natl. Acad. Sci. USA</u>, <u>77</u>: 4030 (1980); and U.S. Pat. Nos. 4,485,045 and 4,544,545. Liposomes with enhanced circulation time are disclosed in U.S. Patent No. 5,013,556.

Particularly useful liposomes can be generated by the reverse-phase evaporation method with a lipid composition comprising phosphatidylcholine, cholesterol, and PEG-derivatized phosphatidylethanolamine (PEG-PE). Liposomes are extruded through filters of defined pore size to yield liposomes with the desired diameter. Fab' fragments of the antibody of the present invention can be conjugated to the liposomes as described in Martin et al., J. Biol. Chem., 257: 286-288 (1982) via a disulfide-interchange reaction. A chemotherapeutic agent (such as Doxorubicin) is optionally contained within the liposome. See Gabizon et al., J. National Cancer Inst., 81(19): 1484 (1989).

M. Pharmaceutical Compositions

The active PRO molecules of the invention (e.g., PRO polypeptides, anti-PRO antibodies, and/or variants of each) as well as other molecules identified by the screening assays disclosed above, can be administered for the treatment of immune related diseases, in the form of pharmaceutical compositions.

Therapeutic formulations of the active PRO molecule, preferably a polypeptide or antibody of the invention, are prepared for storage by mixing the active molecule having the desired degree of purity with optional pharmaceutically acceptable carriers, excipients or stabilizers (Remington's Pharmaceutical Sciences 16th edition, Osol, A. Ed. [1980]), in the form of lyophilized formulations or aqueous solutions. Acceptable carriers, excipients, or stabilizers are nontoxic to recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate, and other organic acids; antioxidants including ascorbic acid and methionine; preservatives (such as octadecyldimethylbenzyl ammonium chloride; hexamethonium chloride; benzalkonium chloride, benzethonium chloride; phenol, butyl or benzyl alcohol; alkyl parabens such as methyl or propyl paraben; catechol; resorcinol; cyclohexanol; 3-pentanol; and mcresol); low molecular weight (less than about 10 residues) polypeptides; proteins, such as serum albumin, gelatin, or immunoglobulins; hydrophilic polymers such as polyvinylpyrrolidone; amino acids such as glycine, glutamine, asparagine, histidine, arginine, or lysine; monosaccharides, disaccharides, and other carbohydrates including glucose, mannose, or dextrins; chelating agents such as EDTA; sugars such as sucrose, mannitol, trehalose or sorbitol; salt-forming counter-ions such as sodium; metal complexes (e.g., Zn-protein complexes); and/or non-ionic surfactants such as TWEENTM, PLURONICSTM or polyethylene glycol (PEG).

Compounds identified by the screening assays disclosed herein can be formulated in an analogous manner, using standard techniques well known in the art.

Lipofections or liposomes can also be used to deliver the PRO molecule into cells. Where antibody fragments are used, the smallest inhibitory fragment which specifically binds to the binding domain of the target protein is preferred. For example, based upon the variable region sequences of an antibody, peptide molecules can be designed which retain the ability to bind the target protein sequence. Such peptides can be synthesized chemically and/or produced by recombinant DNA technology (see, e.g., Marasco et al., Proc. Natl. Acad. Sci. USA 90, 7889-7893 [1993]).

The formulation herein may also contain more than one active compound as necessary for the particular indication being treated, preferably those with complementary activities that do not adversely affect each other. Alternatively, or in addition, the composition may comprise a cytotoxic agent, cytokine or growth inhibitory agent. Such molecules are suitably present in combination in amounts that are effective for the purpose intended.

The active PRO molecules may also be entrapped in microcapsules prepared, for example, by coacervation techniques or by interfacial polymerization, for example, hydroxymethylcellulose or gelatin-microcapsules and poly-(methylmethacylate) microcapsules, respectively, in colloidal drug delivery systems (for example, liposomes, albumin microspheres, microemulsions, nano-particles and nanocapsules) or in macroemulsions. Such techniques are disclosed in *Remington's Pharmaceutical Sciences* 16th edition, Osol, A. Ed. (1980).

The formulations to be used for *in vivo* administration must be sterile. This is readily accomplished by filtration through sterile filtration membranes.

Sustained-release preparations or the PRO molecules may be prepared. Suitable examples of sustained-release preparations include semipermeable matrices of solid hydrophobic polymers containing the antibody, which matrices are in the form of shaped articles, e.g., films, or microcapsules. Examples of sustained-release matrices include polyesters, hydrogels (for example, poly(2-hydroxyethyl-methacrylate), or poly(vinylalcohol)), polylactides (U.S. Pat. No. 3,773,919), copolymers of L-glutamic acid and γ-ethyl-Lglutamate, non-degradable ethylene-vinyl acetate, degradable lactic acid-glycolic acid copolymers such as the LUPRON DEPOTTM (injectable microspheres composed of lactic acid-glycolic acid copolymer and leuprolide acetate), and poly-D-(-)-3-hydroxybutyric acid. While polymers such as ethylene-vinyl acetate and lactic acid-glycolic acid enable release of molecules for over 100 days, certain hydrogels release proteins for shorter time periods. When encapsulated antibodies remain in the body for a long time, they may denature or aggregate as a result of exposure to moisture at 37°C, resulting in a loss of biological activity and possible changes in immunogenicity. Rational strategies can be devised for stabilization depending on the mechanism involved. For example, if the aggregation mechanism is discovered to be intermolecular S-S bond formation through thio-disulfide interchange, stabilization may be achieved by modifying sulfhydryl residues, lyophilizing from acidic solutions, controlling moisture content, using appropriate additives, and developing specific polymer matrix compositions.

N. Methods of Treatment

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It is contemplated that the polypeptides, antibodies and other active compounds of the present invention may be used to treat various immune related diseases and conditions, such as T cell mediated diseases, including those characterized by infiltration of inflammatory cells into a tissue, stimulation of T-

cell proliferation, inhibition of T-cell proliferation, increased or decreased vascular permeability or the inhibition thereof.

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Exemplary conditions or disorders to be treated with the polypeptides, antibodies and other compounds of the invention, include, but are not limited to systemic lupus erythematosis, rheumatoid arthritis, juvenile chronic arthritis, osteoarthritis, spondyloarthropathies, systemic sclerosis (scleroderma), idiopathic inflammatory myopathies (dermatomyositis, polymyositis), Sjögren's syndrome, systemic vasculitis, sarcoidosis, autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria), autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia), thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis), diabetes mellitus, immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis), demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy, hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis, inflammatory bowel disease (ulcerative colitis: Crohn's disease), gluten-sensitive enteropathy, and Whipple's disease, autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis, allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria, immunologic diseases of the lung such as eosinophilic pneumonias, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis, transplantation associated diseases including graft rejection and graft -versus-host-disease.

In systemic lupus erythematosus, the central mediator of disease is the production of auto-reactive antibodies to self proteins/tissues and the subsequent generation of immune-mediated inflammation. Antibodies either directly or indirectly mediate tissue injury. Though T lymphocytes have not been shown to be directly involved in tissue damage, T lymphocytes are required for the development of auto-reactive antibodies. The genesis of the disease is thus T lymphocyte dependent. Multiple organs and systems are affected clinically including kidney, lung, musculoskeletal system, mucocutaneous, eye, central nervous system, cardiovascular system, gastrointestinal tract, bone marrow and blood.

Rheumatoid arthritis (RA) is a chronic systemic autoimmune inflammatory disease that mainly involves the synovial membrane of multiple joints with resultant injury to the articular cartilage. The pathogenesis is T lymphocyte dependent and is associated with the production of rheumatoid factors, auto-antibodies directed against self IgG, with the resultant formation of immune complexes that attain high levels in joint fluid and blood. These complexes in the joint may induce the marked infiltrate of lymphocytes and monocytes into the synovium and subsequent marked synovial changes; the joint space/fluid if infiltrated by similar cells with the addition of numerous neutrophils. Tissues affected are primarily the joints, often in symmetrical pattern. However, extra-articular disease also occurs in two major forms. One form is the development of extra-articular lesions with ongoing progressive joint disease and typical lesions of pulmonary fibrosis, vasculitis, and cutaneous ulcers. The second form of extra-articular disease is the so called Felty's syndrome which occurs late in the RA disease course, sometimes after joint disease has become quiescent, and involves the presence of neutropenia, thrombocytopenia and splenomegaly. This can be accompanied by vasculitis in multiple organs with formations of infarcts, skin

ulcers and gangrene. Patients often also develop rheumatoid nodules in the subcutis tissue overlying affected joints; the nodules late stage have necrotic centers surrounded by a mixed inflammatory cell infiltrate. Other manifestations which can occur in RA include: pericarditis, pleuritis, coronary arteritis, intestitial pneumonitis with pulmonary fibrosis, keratoconjunctivitis sicca, and rhematoid nodules.

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Juvenile chronic arthritis is a chronic idiopathic inflammatory disease which begins often at less than 16 years of age. Its phenotype has some similarities to RA; some patients which are rhematoid factor positive are classified as juvenile rheumatoid arthritis. The disease is sub-classified into three major categories: pauciarticular, polyarticular, and systemic. The arthritis can be severe and is typically destructive and leads to joint ankylosis and retarded growth. Other manifestations can include chronic anterior uveitis and systemic amyloidosis.

Spondyloarthropathies are a group of disorders with some common clinical features and the common association with the expression of HLA-B27 gene product. The disorders include: ankylosing sponylitis, Reiter's syndrome (reactive arthritis), arthritis associated with inflammatory bowel disease, spondylitis associated with psoriasis, juvenile onset spondyloarthropathy and undifferentiated spondyloarthropathy. Distinguishing features include sacroileitis with or without spondylitis; inflammatory asymmetric arthritis; association with HLA-B27 (a serologically defined allele of the HLA-B locus of class I MHC); ocular inflammation, and absence of autoantibodies associated with other rheumatoid disease. The cell most implicated as key to induction of the disease is the CD8+ T lymphocyte, a cell which targets antigen presented by class I MHC molecules. CD8+ T cells may react against the class I MHC allele HLA-B27 as if it were a foreign peptide expressed by MHC class I molecules. It has been hypothesized that an epitope of HLA-B27 may mimic a bacterial or other microbial antigenic epitope and thus induce a CD8+ T cells response.

Systemic sclerosis (scleroderma) has an unknown etiology. A hallmark of the disease is induration of the skin; likely this is induced by an active inflammatory process. Scleroderma can be localized or systemic; vascular lesions are common and endothelial cell injury in the microvasculature is an early and important event in the development of systemic sclerosis; the vascular injury may be immune mediated. An immunologic basis is implied by the presence of mononuclear cell infiltrates in the cutaneous lesions and the presence of anti-nuclear antibodies in many patients. ICAM-1 is often upregulated on the cell surface of fibroblasts in skin lesions suggesting that T cell interaction with these cells may have a role in the pathogenesis of the disease. Other organs involved include: the gastrointestinal tract: smooth muscle atrophy and fibrosis resulting in abnormal peristalsis/motility; kidney: concentric subendothelial intimal proliferation affecting small arcuate and interlobular arteries with resultant reduced renal cortical blood flow, results in proteinuria, azotemia and hypertension; skeletal muscle: atrophy, interstitial fibrosis; inflammation; lung: interstitial pneumonitis and interstitial fibrosis; and heart: contraction band necrosis, scarring/fibrosis.

Idiopathic inflammatory myopathies including dermatomyositis, polymyositis and others are disorders of chronic muscle inflammation of unknown etiology resulting in muscle weakness. Muscle injury/inflammation is often symmetric and progressive. Autoantibodies are associated with most forms. These myositis-specific autoantibodies are directed against and inhibit the function of components, proteins and RNA's, involved in protein synthesis.

Sjögren's syndrome is due to immune-mediated inflammation and subsequent functional destruction of the tear glands and salivary glands. The disease can be associated with or accompanied by inflammatory connective tissue diseases. The disease is associated with autoantibody production against Ro and La antigens, both of which are small RNA-protein complexes. Lesions result in keratoconjunctivitis sicca, xerostomia, with other manifestations or associations including bilary cirrhosis, peripheral or sensory neuropathy, and palpable purpura.

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Systemic vasculitis are diseases in which the primary lesion is inflammation and subsequent damage to blood vessels which results in ischemia/necrosis/degeneration to tissues supplied by the affected vessels and eventual end-organ dysfunction in some cases. Vasculitides can also occur as a secondary lesion or sequelae to other immune-inflammatory mediated diseases such as rheumatoid arthritis, systemic sclerosis, etc., particularly in diseases also associated with the formation of immune complexes. Diseases in the primary systemic vasculitis group include: systemic necrotizing vasculitis: polyarteritis nodosa, allergic angiitis and granulomatosis, polyangiitis; Wegener's granulomatosis; lymphomatoid granulomatosis; and giant cell arteritis. Miscellaneous vasculitides include: mucocutaneous lymph node syndrome (MLNS or Kawasaki's disease), isolated CNS vasculitis, Behet's disease, thromboangiitis obliterans (Buerger's disease) and cutaneous necrotizing venulitis. The pathogenic mechanism of most of the types of vasculitis listed is believed to be primarily due to the deposition of immunoglobulin complexes in the vessel wall and subsequent induction of an inflammatory response either via ADCC, complement activation, or both.

Sarcoidosis is a condition of unknown etiology which is characterized by the presence of epithelioid granulomas in nearly any tissue in the body; involvement of the lung is most common. The pathogenesis involves the persistence of activated macrophages and lymphoid cells at sites of the disease with subsequent chronic sequelae resultant from the release of locally and systemically active products released by these cell types.

Autoimmune hemolytic anemia including autoimmune hemolytic anemia, immune pancytopenia, and paroxysmal noctural hemoglobinuria is a result of production of antibodies that react with antigens expressed on the surface of red blood cells (and in some cases other blood cells including platelets as well) and is a reflection of the removal of those antibody coated cells via complement mediated lysis and/or ADCC/Fc-receptor-mediated mechanisms.

In autoimmune thrombocytopenia including thrombocytopenic purpura, and immune-mediated thrombocytopenia in other clinical settings, platelet destruction/removal occurs as a result of either antibody or complement attaching to platelets and subsequent removal by complement lysis, ADCC or FC-receptor mediated mechanisms.

Thyroiditis including Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, and atrophic thyroiditis, are the result of an autoimmune response against thyroid antigens with production of antibodies that react with proteins present in and often specific for the thyroid gland. Experimental models exist including spontaneous models: rats (BUF and BB rats) and chickens (obese chicken strain); inducible models: immunization of animals with either thyroglobulin, thyroid microsomal antigen (thyroid peroxidase).

Type I diabetes mellitus or insulin-dependent diabetes is the autoimmune destruction of pancreatic islet β cells; this destruction is mediated by auto-antibodies and auto-reactive T cells. Antibodies to insulin or the insulin receptor can also produce the phenotype of insulin-non-responsiveness.

Immune mediated renal diseases, including glomerulonephritis and tubulointerstitial nephritis, are the result of antibody or T lymphocyte mediated injury to renal tissue either directly as a result of the production of autoreactive antibodies or T cells against renal antigens or indirectly as a result of the deposition of antibodies and/or immune complexes in the kidney that are reactive against other, non-renal antigens. Thus other immune-mediated diseases that result in the formation of immune-complexes can also induce immune mediated renal disease as an indirect sequelae. Both direct and indirect immune mechanisms result in inflammatory response that produces/induces lesion development in renal tissues with resultant organ function impairment and in some cases progression to renal failure. Both humoral and cellular immune mechanisms can be involved in the pathogenesis of lesions.

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Demyelinating diseases of the central and peripheral nervous systems, including Multiple Sclerosis; idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome; and Chronic Inflammatory Demyelinating Polyneuropathy, are believed to have an autoimmune basis and result in nerve demyelination as a result of damage caused to oligodendrocytes or to myelin directly. In MS there is evidence to suggest that disease induction and progression is dependent on T lymphocytes. Multiple Sclerosis is a demyelinating disease that is T lymphocyte-dependent and has either a relapsing-remitting course or a chronic progressive course. The etiology is unknown; however, viral infections, genetic predisposition, environment, and autoimmunity all contribute. Lesions contain infiltrates of predominantly T lymphocyte mediated, microglial cells and infiltrating macrophages; CD4+ T lymphocytes are the predominant cell type at lesions. The mechanism of oligodendrocyte cell death and subsequent demyelination is not known but is likely T lymphocyte driven.

Inflammatory and Fibrotic Lung Disease, including Eosinophilic Pneumonias; Idiopathic Pulmonary Fibrosis, and Hypersensitivity Pneumonitis may involve a disregulated immune-inflammatory response. Inhibition of that response would be of therapeutic benefit.

Autoimmune or Immune-mediated Skin Disease including Bullous Skin Diseases, Erythema Multiforme, and Contact Dermatitis are mediated by auto-antibodies, the genesis of which is T lymphocyte-dependent.

Psoriasis is a T lymphocyte-mediated inflammatory disease. Lesions contain infiltrates of T lymphocytes, macrophages and antigen processing cells, and some neutrophils.

Allergic diseases, including asthma; allergic rhinitis; atopic dermatitis; food hypersensitivity; and urticaria are T lymphocyte dependent. These diseases are predominantly mediated by T lymphocyte induced inflammation, IgE mediated-inflammation or a combination of both.

Transplantation associated diseases, including Graft rejection and Graft-Versus-Host-Disease (GVHD) are T lymphocyte-dependent; inhibition of T lymphocyte function is ameliorative.

Other diseases in which intervention of the immune and/or inflammatory response have benefit are infectious disease including but not limited to viral infection (including but not limited to AIDS, hepatitis A, B, C, D, E and herpes) bacterial infection, fungal infections, and protozoal and parasitic infections (molecules (or derivatives/agonists) which stimulate the MLR can be utilized therapeutically to enhance the

immune response to infectious agents), diseases of immunodeficiency (molecules/derivatives/agonists) which stimulate the MLR can be utilized therapeutically to enhance the immune response for conditions of inherited, acquired, infectious induced (as in HIV infection), or introgenic (i.e., as from chemotherapy) immunodeficiency, and neoplasia.

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It has been demonstrated that some human cancer patients develop an antibody and/or T lymphocyte response to antigens on neoplastic cells. It has also been shown in animal models of neoplasia that enhancement of the immune response can result in rejection or regression of that particular neoplasm. Molecules that enhance the T lymphocyte response in the MLR have utility in vivo in enhancing the immune response against neoplasia. Molecules which enhance the T lymphocyte proliferative response in the MLR (or small molecule agonists or antibodies that affected the same receptor in an agonistic fashion) can be used therapeutically to treat cancer. Molecules that inhibit the lymphocyte response in the MLR also function in vivo during neoplasia to suppress the immune response to a neoplasm; such molecules can either be expressed by the neoplastic cells themselves or their expression can be induced by the neoplasm in other cells. Antagonism of such inhibitory molecules (either with antibody, small molecule antagonists or other means) enhances immune-mediated tumor rejection.

Additionally, inhibition of molecules with proinflammatory properties may have therapeutic benefit in reperfusion injury; stroke; myocardial infarction; atherosclerosis; acute lung injury; hemorrhagic shock; burn; sepsis/septic shock; acute tubular necrosis; endometriosis; degenerative joint disease and pancreatis.

The compounds of the present invention, e.g., polypeptides or antibodies, are administered to a mammal, preferably a human, in accord with known methods, such as intravenous administration as a bolus or by continuous infusion over a period of time, by intramuscular, intraperitoneal, intracerobrospinal, subcutaneous, intra-articular, intrasynovial, intrathecal, oral, topical, or inhalation (intranasal, intrapulmonary) routes. Intravenous or inhaled administration of polypeptides and antibodies is preferred.

In immunoadjuvant therapy, other therapeutic regimens, such administration of an anti-cancer agent, may be combined with the administration of the proteins, antibodies or compounds of the instant invention. For example, the patient to be treated with a the immunoadjuvant of the invention may also receive an anti-cancer agent (chemotherapeutic agent) or radiation therapy. Preparation and dosing schedules for such chemotherapeutic agents may be used according to manufacturers' instructions or as determined empirically by the skilled practitioner. Preparation and dosing schedules for such chemotherapy are also described in *Chemotherapy Service* Ed., M.C. Perry, Williams & Wilkins, Baltimore, MD (1992). The chemotherapeutic agent may precede, or follow administration of the immunoadjuvant or may be given simultaneously therewith. Additionally, an anti-estrogen compound such as tamoxifen or an anti-progesterone such as onapristone (see, EP 616812) may be given in dosages known for such molecules.

It may be desirable to also administer antibodies against other immune disease associated or tumor associated antigens, such as antibodies which bind to CD20, CD11a, CD18, ErbB2, EGFR, ErbB3, ErbB4, or vascular endothelial factor (VEGF). Alternatively, or in addition, two or more antibodies binding the same or two or more different antigens disclosed herein may be coadministered to the patient. Sometimes, it may be beneficial to also administer one or more cytokines to the patient. In one embodiment, the PRO polypeptides are coadministered with a growth inhibitory agent. For example, the growth inhibitory agent may be administered first, followed by a PRO polypeptide. However, simultaneous administration or

administration first is also contemplated. Suitable dosages for the growth inhibitory agent are those presently used and may be lowered due to the combined action (synergy) of the growth inhibitory agent and the PRO polypeptide.

For the treatment or reduction in the severity of immune related disease, the appropriate dosage of an a compound of the invention will depend on the type of disease to be treated, as defined above, the severity and course of the disease, whether the agent is administered for preventive or therapeutic purposes, previous therapy, the patient's clinical history and response to the compound, and the discretion of the attending physician. The compound is suitably administered to the patient at one time or over a series of treatments.

For example, depending on the type and severity of the disease, about 1 μ g/kg to 15 mg/kg (e.g., 0.1-20 mg/kg) of polypeptide or antibody is an initial candidate dosage for administration to the patient, whether, for example, by one or more separate administrations, or by continuous infusion. A typical daily dosage might range from about 1 μ g/kg to 100 mg/kg or more, depending on the factors mentioned above. For repeated administrations over several days or longer, depending on the condition, the treatment is sustained until a desired suppression of disease symptoms occurs. However, other dosage regimens may be useful. The progress of this therapy is easily monitored by conventional techniques and assays.

O. Articles of Manufacture

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In another embodiment of the invention, an article of manufacture containing materials (e.g., comprising a PRO molecule) useful for the diagnosis or treatment of the disorders described above is provided. The article of manufacture comprises a container and an instruction. Suitable containers include, for example, bottles, vials, syringes, and test tubes. The containers may be formed from a variety of materials such as glass or plastic. The container holds a composition which is effective for diagnosing or treating the condition and may have a sterile access port (for example the container may be an intravenous solution bag or a vial having a stopper pierceable by a hypodermic injection needle). The active agent in the composition is usually a polypeptide or an antibody of the invention. An instruction or label on, or associated with, the container indicates that the composition is used for diagnosing or treating the condition of choice. The article of manufacture may further comprise a second container comprising a pharmaceutically-acceptable buffer, such as phosphate-buffered saline, Ringer's solution and dextrose solution. It may further include other materials desirable from a commercial and user standpoint, including other buffers, diluents, filters, needles, syringes, and package inserts with instructions for use.

P. <u>Diagnosis and Prognosis of Immune Related Disease</u>

Cell surface proteins, such as proteins which are overexpressed in certain immune related diseases, are excellent targets for drug candidates or disease treatment. The same proteins along with secreted proteins encoded by the genes amplified in immune related disease states find additional use in the diagnosis and prognosis of these diseases. For example, antibodies directed against the protein products of genes amplified in multiple sclerosis, rheumatoid arthritis, or another immune related disease, can be used as diagnostics or prognostics.

For example, antibodies, including antibody fragments, can be used to qualitatively or quantitatively detect the expression of proteins encoded by amplified or overexpressed genes ("marker gene products"). The antibody preferably is equipped with a detectable, e.g., fluorescent label, and binding can be

monitored by light microscopy, flow cytometry, fluorimetry, or other techniques known in the art. These techniques are particularly suitable, if the overexpressed gene encodes a cell surface protein Such binding assays are performed essentially as described above.

In situ detection of antibody binding to the marker gene products can be performed, for example, by immunofluorescence or immunoelectron microscopy. For this purpose, a histological specimen is removed from the patient, and a labeled antibody is applied to it, preferably by overlaying the antibody on a biological sample. This procedure also allows for determining the distribution of the marker gene product in the tissue examined. It will be apparent for those skilled in the art that a wide variety of histological methods are readily available for in situ detection.

The following examples are offered for illustrative purposes only, and are not intended to limit the scope of the present invention in any way.

All patent and literature references cited in the present specification are hereby incorporated by reference in their entirety.

EXAMPLES

Commercially available reagents referred to in the examples were used according to manufacturer's instructions unless otherwise indicated. The source of those cells identified in the following examples, and throughout the specification, by ATCC accession numbers is the American Type Culture Collection, Manassas, VA.

EXAMPLE 1: Microarray analysis of stimulated T-cells

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Nucleic acid microarrays, often containing thousands of gene sequences, are useful for identifying differentially expressed genes in diseased tissues as compared to their normal counterparts. Using nucleic acid microarrays, test and control mRNA samples from test and control tissue samples are reverse transcribed and labeled to generate cDNA probes. The cDNA probes are then hybridized to an array of nucleic acids immobilized on a solid support. The array is configured such that the sequence and position of each member of the array is known. For example, a selection of genes known to be expressed in certain disease states may be arrayed on a solid support. Hybridization of a labeled probe with a particular array member indicates that the sample from which the probe was derived expresses that gene. If the hybridization signal of a probe from a test (in this instance, activated CD4+ T cells) sample is greater than hybridization signal of a probe from a control (in this instance, non-stimulated CD4 + T cells) sample, the gene or genes overexpressed in the test tissue are identified. The implication of this result is that an overexpressed protein in a test tissue is useful not only as a diagnostic marker for the presence of the disease condition, but also as a therapeutic target for treatment of the disease condition.

The methodology of hybridization of nucleic acids and microarray technology is well known in the art. In one example, the specific preparation of nucleic acids for hybridization and probes, slides, and hybridization conditions are all detailed in PCT Patent Application Serial No. PCT/US01/10482, filed on March 30, 2001 and which is herein incorporated by reference.

In this experiment, CD4+ T cells were purified from a single donor using the RossetteSep™ protocol from (Stem Cell Technologies, Vancouver BC) which contains anti-CD8, anti-CD16, anti-CD19, anti-CD36 and anti-CD56 antibodies used to produce a population of isolated CD4+ T cells. Isolated CD4+

T cells were activated with an anti-CD3 antibody (used at a concentration that does not stimulate proliferation) together with either ICAM-1 or anti-CD28 antibody. At 24 or 72 hours cells were harvested, RNA extracted and analysis run on Affimax (Affymetrix Inc. Santa Clara, CA) microarray chips. Non-stimulated (resting) cells were harvested immediately after purification, and subjected to the same analysis. Genes were compared whose expression was upregulated at either of the two timepoints in activated vs. resting cells.

Below are the results of these experiments, demonstrating that various PRO polypeptides of the present invention are differentially expressed in isolated CD4 + T cells activated by anti-CD3/ICAM-1 or anti-CD3/anti-CD28 as compared to isolated resting CD4+ T cells. As described above, these data demonstrate that the PRO polypeptides of the present invention are useful not only as diagnostic markers for the presence of one or more immune disorders, but also serve as therapeutic targets for the treatment of those immune disorders.

The results of this experient are Figures 1-7589 show increase or decrease in expression upon stimulation with anti-CD3/ICAM1 and also show increase or decrease in expression upon stimulation with anti-CD3/anti-CD28. The nucleic acids and encoded proteins of Figure 946, Figure 1520, Figure 1574, Figure 1622, Figure 1816, Figure 2433, Figure 2986, Figure 3220, Figure 4120 and Figure 5421 are significantly overexpressed in isolated CD4 + T cells activated by anti-CD3/ICAM-1 or anti-CD3/anti-CD28 as compared to isolated resting CD4+ T cells.

EXAMPLE 2: Use of PRO as a hybridization probe

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The following method describes use of a nucleotide sequence encoding PRO as a hybridization probe.

DNA comprising the coding sequence of full-length or mature PRO as disclosed herein is employed as a probe to screen for homologous DNAs (such as those encoding naturally-occurring variants of PRO) in human tissue cDNA libraries or human tissue genomic libraries.

Hybridization and washing of filters containing either library DNAs is performed under the following high stringency conditions. Hybridization of radiolabeled PRO-derived probe to the filters is performed in a solution of 50% formamide, 5x SSC, 0.1% SDS, 0.1% sodium pyrophosphate, 50 mM sodium phosphate, pH 6.8, 2x Denhardt's solution, and 10% dextran sulfate at 42°C for 20 hours. Washing of the filters is performed in an aqueous solution of 0.1x SSC and 0.1% SDS at 42°C.

DNAs having a desired sequence identity with the DNA encoding full-length native sequence PRO can then be identified using standard techniques known in the art.

EXAMPLE 3: Expression of PRO in E. coli

This example illustrates preparation of an unglycosylated form of PRO by recombinant expression in *E. coli*.

The DNA sequence encoding PRO is initially amplified using selected PCR primers. The primers should contain restriction enzyme sites which correspond to the restriction enzyme sites on the selected expression vector. A variety of expression vectors may be employed. An example of a suitable vector is pBR322 (derived from *E. coli*; see Bolivar et al., <u>Gene</u>, <u>2</u>:95 (1977)) which contains genes for ampicillin and

tetracycline resistance. The vector is digested with restriction enzyme and dephosphorylated. The PCR amplified sequences are then ligated into the vector. The vector will preferably include sequences which encode for an antibiotic resistance gene, a trp promoter, a polyhis leader (including the first six STII codons, polyhis sequence, and enterokinase cleavage site), the PRO coding region, lambda transcriptional terminator, and an argU gene.

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The ligation mixture is then used to transform a selected *E. coli* strain using the methods described in Sambrook et al., <u>supra</u>. Transformants are identified by their ability to grow on LB plates and antibiotic resistant colonies are then selected. Plasmid DNA can be isolated and confirmed by restriction analysis and DNA sequencing.

Selected clones can be grown overnight in liquid culture medium such as LB broth supplemented with antibiotics. The overnight culture may subsequently be used to inoculate a larger scale culture. The cells are then grown to a desired optical density, during which the expression promoter is turned on.

After culturing the cells for several more hours, the cells can be harvested by centrifugation. The cell pellet obtained by the centrifugation can be solubilized using various agents known in the art, and the solubilized PRO protein can then be purified using a metal chelating column under conditions that allow tight binding of the protein.

PRO may be expressed in *E. coli* in a poly-His tagged form, using the following procedure. The DNA encoding PRO is initially amplified using selected PCR primers. The primers will contain restriction enzyme sites which correspond to the restriction enzyme sites on the selected expression vector, and other useful sequences providing for efficient and reliable translation initiation, rapid purification on a metal chelation column, and proteolytic removal with enterokinase. The PCR-amplified, poly-His tagged sequences are then ligated into an expression vector, which is used to transform an *E. coli* host based on strain 52 (W3110 fuhA(tonA) lon galE rpoHts(htpRts) clpP(lacIq). Transformants are first grown in LB containing 50 mg/ml carbenicillin at 30°C with shaking until an O.D.600 of 3-5 is reached. Cultures are then diluted 50-100 fold into CRAP media (prepared by mixing 3.57 g (NH₄)₂SO₄, 0.71 g sodium citrate•2H2O, 1.07 g KCl, 5.36 g Difco yeast extract, 5.36 g Sheffield hycase SF in 500 mL water, as well as 110 mM MPOS, pH 7.3, 0.55% (w/v) glucose and 7 mM MgSO₄) and grown for approximately 20-30 hours at 30°C with shaking. Samples are removed to verify expression by SDS-PAGE analysis, and the bulk culture is centrifuged to pellet the cells. Cell pellets are frozen until purification and refolding.

E. coli paste from 0.5 to 1 L fermentations (6-10 g pellets) is resuspended in 10 volumes (w/v) in 7 M guanidine, 20 mM Tris, pH 8 buffer. Solid sodium sulfite and sodium tetrathionate is added to make final concentrations of 0.1M and 0.02 M, respectively, and the solution is stirred overnight at 4°C. This step results in a denatured protein with all cysteine residues blocked by sulfitolization. The solution is centrifuged at 40,000 rpm in a Beckman Ultracentifuge for 30 min. The supernatant is diluted with 3-5 volumes of metal chelate column buffer (6 M guanidine, 20 mM Tris, pH 7.4) and filtered through 0.22 micron filters to clarify. The clarified extract is loaded onto a 5 ml Qiagen Ni-NTA metal chelate column equilibrated in the metal chelate column buffer. The column is washed with additional buffer containing 50 mM imidazole (Calbiochem, Utrol grade), pH 7.4. The protein is eluted with buffer containing 250 mM imidazole. Fractions containing the desired protein are pooled and stored at 4°C. Protein concentration is estimated by its absorbance at 280 nm using the calculated extinction coefficient based on its amino acid sequence.

The proteins are refolded by diluting the sample slowly into freshly prepared refolding buffer consisting of: 20 mM Tris, pH 8.6, 0.3 M NaCl, 2.5 M urea, 5 mM cysteine, 20 mM glycine and 1 mM EDTA. Refolding volumes are chosen so that the final protein concentration is between 50 to 100 micrograms/ml. The refolding solution is stirred gently at 4°C for 12-36 hours. The refolding reaction is quenched by the addition of TFA to a final concentration of 0.4% (pH of approximately 3). Before further purification of the protein, the solution is filtered through a 0.22 micron filter and acetonitrile is added to 2-10% final concentration. The refolded protein is chromatographed on a Poros R1/H reversed phase column using a mobile buffer of 0.1% TFA with elution with a gradient of acetonitrile from 10 to 80%. Aliquots of fractions with A280 absorbance are analyzed on SDS polyacrylamide gels and fractions containing homogeneous refolded protein are pooled. Generally, the properly refolded species of most proteins are eluted at the lowest concentrations of acetonitrile since those species are the most compact with their hydrophobic interiors shielded from interaction with the reversed phase resin. Aggregated species are usually eluted at higher acetonitrile concentrations. In addition to resolving misfolded forms of proteins from the desired form, the reversed phase step also removes endotoxin from the samples.

Fractions containing the desired folded PRO polypeptide are pooled and the acetonitrile removed using a gentle stream of nitrogen directed at the solution. Proteins are formulated into 20 mM Hepes, pH 6.8 with 0.14 M sodium chloride and 4% mannitol by dialysis or by gel filtration using G25 Superfine (Pharmacia) resins equilibrated in the formulation buffer and sterile filtered.

Many of the PRO polypeptides disclosed herein were successfully expressed as described above.

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EXAMPLE 4: Expression of PRO in mammalian cells

This example illustrates preparation of a potentially glycosylated form of PRO by recombinant expression in mammalian cells.

The vector, pRK5 (see EP 307,247, published March 15, 1989), is employed as the expression vector. Optionally, the PRO DNA is ligated into pRK5 with selected restriction enzymes to allow insertion of the PRO DNA using ligation methods such as described in Sambrook et al., <u>supra</u>. The resulting vector is called pRK5-PRO.

In one embodiment, the selected host cells may be 293 cells. Human 293 cells (ATCC CCL 1573) are grown to confluence in tissue culture plates in medium such as DMEM supplemented with fetal calf serum and optionally, nutrient components and/or antibiotics. About 10 µg pRK5-PRO DNA is mixed with about 1 µg DNA encoding the VA RNA gene [Thimmappaya et al., Cell, 31:543 (1982)] and dissolved in 500 µl of 1 mM Tris-HCl, 0.1 mM EDTA, 0.227 M CaCl₂. To this mixture is added, dropwise, 500 µl of 50 mM HEPES (pH 7.35), 280 mM NaCl, 1.5 mM NaPO₄, and a precipitate is allowed to form for 10 minutes at 25°C. The precipitate is suspended and added to the 293 cells and allowed to settle for about four hours at 37°C. The culture medium is aspirated off and 2 ml of 20% glycerol in PBS is added for 30 seconds. The 293 cells are then washed with serum free medium, fresh medium is added and the cells are incubated for about 5 days.

Approximately 24 hours after the transfections, the culture medium is removed and replaced with culture medium (alone) or culture medium containing 200 μ Ci/ml ³⁵S-cysteine and 200 μ Ci/ml ³⁵S-methionine. After a 12 hour incubation, the conditioned medium is collected, concentrated on a spin filter.

and loaded onto a 15% SDS gel. The processed gel may be dried and exposed to film for a selected period of time to reveal the presence of PRO polypeptide. The cultures containing transfected cells may undergo further incubation (in serum free medium) and the medium is tested in selected bioassays.

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In an alternative technique, PRO may be introduced into 293 cells transiently using the dextran sulfate method described by Somparyrac et al., Proc. Natl. Acad. Sci., 12:7575 (1981). 293 cells are grown to maximal density in a spinner flask and 700 μ g pRK5-PRO DNA is added. The cells are first concentrated from the spinner flask by centrifugation and washed with PBS. The DNA-dextran precipitate is incubated on the cell pellet for four hours. The cells are treated with 20% glycerol for 90 seconds, washed with tissue culture medium, and re-introduced into the spinner flask containing tissue culture medium, 5 μ g/ml bovine insulin and 0.1 μ g/ml bovine transferrin. After about four days, the conditioned media is centrifuged and filtered to remove cells and debris. The sample containing expressed PRO can then be concentrated and purified by any selected method, such as dialysis and/or column chromatography.

In another embodiment, PRO can be expressed in CHO cells. The pRK5-PRO can be transfected into CHO cells using known reagents such as CaPO₄ or DEAE-dextran. As described above, the cell cultures can be incubated, and the medium replaced with culture medium (alone) or medium containing a radiolabel such as ³⁵S-methionine. After determining the presence of PRO polypeptide, the culture medium may be replaced with serum free medium. Preferably, the cultures are incubated for about 6 days, and then the conditioned medium is harvested. The medium containing the expressed PRO can then be concentrated and purified by any selected method.

Epitope-tagged PRO may also be expressed in host CHO cells. The PRO may be subcloned out of the pRK5 vector. The subclone insert can undergo PCR to fuse in frame with a selected epitope tag such as a poly-his tag into a Baculovirus expression vector. The poly-his tagged PRO insert can then be subcloned into a SV40 promoter/enhancer containing vector containing a selection marker such as DHFR for selection of stable clones. Finally, the CHO cells can be transfected (as described above) with the SV40 promoter/enhancer containing vector. Labeling may be performed, as described above, to verify expression. The culture medium containing the expressed poly-His tagged PRO can then be concentrated and purified by any selected method, such as by Ni²⁺-chelate affinity chromatography.

PRO may also be expressed in CHO and/or COS cells by a transient expression procedure or in CHO cells by another stable expression procedure.

Stable expression in CHO cells is performed using the following procedure. The proteins are expressed as an IgG construct (immunoadhesin), in which the coding sequences for the soluble forms (e.g. extracellular domains) of the respective proteins are fused to an IgG1 constant region sequence containing the hinge, CH2 and CH2 domains and/or is a poly-His tagged form.

Following PCR amplification, the respective DNAs are subcloned in a CHO expression vector using standard techniques as described in Ausubel et al., <u>Current Protocols of Molecular Biology</u>, Unit 3.16, John Wiley and Sons (1997). CHO expression vectors are constructed to have compatible restriction sites 5' and 3' of the DNA of interest to allow the convenient shuttling of cDNA's. The vector used expression in CHO cells is as described in Lucas et al., <u>Nucl. Acids Res.</u> 24:9 (1774-1779 (1996), and uses the SV40 early promoter/enhancer to drive expression of the cDNA of interest and dihydrofolate reductase (DHFR). DHFR expression permits selection for stable maintenance of the plasmid following transfection.

Twelve micrograms of the desired plasmid DNA is introduced into approximately 10 million CHO cells using commercially available transfection reagents Superfect[®] (Quiagen), Dosper[®] or Fugene[®] (Boehringer Mannheim). The cells are grown as described in Lucas et al., <u>supra</u>. Approximately 3 x 10⁻⁷ cells are frozen in an ampule for further growth and production as described below.

The ampules containing the plasmid DNA are thawed by placement into water bath and mixed by vortexing. The contents are pipetted into a centrifuge tube containing 10 mL of media and centrifuged at 1000 rpm for 5 minutes. The supernatant is aspirated and the cells are resuspended in 10 mL of selective media (0.2 µm filtered PS20 with 5% 0.2 µm diafiltered fetal bovine serum). The cells are then aliquoted into a 100 mL spinner containing 90 mL of selective media. After 1-2 days, the cells are transferred into a 250 mL spinner filled with 150 mL selective growth medium and incubated at 37°C. After another 2-3 days, 250 mL, 500 mL and 2000 mL spinners are seeded with 3 x 105 cells/mL. The cell media is exchanged with fresh media by centrifugation and resuspension in production medium. Although any suitable CHO media may be employed, a production medium described in U.S. Patent No. 5,122,469, issued June 16, 1992 may actually be used. A 3L production spinner is seeded at 1.2 x 10⁶ cells/mL. On day 0, pH is determined. On day 1, the spinner is sampled and sparging with filtered air is commenced. On day 2, the spinner is sampled, the temperature shifted to 33°C, and 30 mL of 500 g/L glucose and 0.6 mL of 10% antifoam (e.g., 35%) polydimethylsiloxane emulsion, Dow Corning 365 Medical Grade Emulsion) taken. Throughout the production, the pH is adjusted as necessary to keep it at around 7.2. After 10 days, or until the viability dropped below 70%, the cell culture is harvested by centrifugation and filtering through a 0.22 µm filter. The filtrate was either stored at 4°C or immediately loaded onto columns for purification.

For the poly-His tagged constructs, the proteins are purified using a Ni-NTA column (Qiagen). Before purification, imidazole is added to the conditioned media to a concentration of 5 mM. The conditioned media is pumped onto a 6 ml Ni-NTA column equilibrated in 20 mM Hepes, pH 7.4, buffer containing 0.3 M NaCl and 5 mM imidazole at a flow rate of 4-5 ml/min. at 4°C. After loading, the column is washed with additional equilibration buffer and the protein eluted with equilibration buffer containing 0.25 M imidazole. The highly purified protein is subsequently desalted into a storage buffer containing 10 mM Hepes, 0.14 M NaCl and 4% mannitol, pH 6.8, with a 25 ml G25 Superfine (Pharmacia) column and stored at -80°C.

Immunoadhesin (Fc-containing) constructs are purified from the conditioned media as follows. The conditioned medium is pumped onto a 5 ml Protein A column (Pharmacia) which had been equilibrated in 20 mM Na phosphate buffer, pH 6.8. After loading, the column is washed extensively with equilibration buffer before elution with 100 mM citric acid, pH 3.5. The eluted protein is immediately neutralized by collecting 1 ml fractions into tubes containing 275 µl of 1 M Tris buffer, pH 9. The highly purified protein is subsequently desalted into storage buffer as described above for the poly-His tagged proteins. The homogeneity is assessed by SDS polyacrylamide gels and by N-terminal amino acid sequencing by Edman degradation.

Many of the PRO polypeptides disclosed herein were successfully expressed as described above.

EXAMPLE 5: Expression of PRO in Yeast

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The following method describes recombinant expression of PRO in yeast.

First, yeast expression vectors are constructed for intracellular production or secretion of PRO from the ADH2/GAPDH promoter. DNA encoding PRO and the promoter is inserted into suitable restriction enzyme sites in the selected plasmid to direct intracellular expression of PRO. For secretion, DNA encoding PRO can be cloned into the selected plasmid, together with DNA encoding the ADH2/GAPDH promoter, a native PRO signal peptide or other mammalian signal peptide, or, for example, a yeast alpha-factor or invertase secretory signal/leader sequence, and linker sequences (if needed) for expression of PRO.

Yeast cells, such as yeast strain AB110, can then be transformed with the expression plasmids described above and cultured in selected fermentation media. The transformed yeast supernatants can be analyzed by precipitation with 10% trichloroacetic acid and separation by SDS-PAGE, followed by staining of the gels with Coomassie Blue stain.

Recombinant PRO can subsequently be isolated and purified by removing the yeast cells from the fermentation medium by centrifugation and then concentrating the medium using selected cartridge filters. The concentrate containing PRO may further be purified using selected column chromatography resins.

Many of the PRO polypeptides disclosed herein were successfully expressed as described above.

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EXAMPLE 6: Expression of PRO in Baculovirus-Infected Insect Cells

The following method describes recombinant expression of PRO in Baculovirus-infected insect cells.

The sequence coding for PRO is fused upstream of an epitope tag contained within a baculovirus expression vector. Such epitope tags include poly-his tags and immunoglobulin tags (like Fc regions of IgG). A variety of plasmids may be employed, including plasmids derived from commercially available plasmids such as pVL1393 (Novagen). Briefly, the sequence encoding PRO or the desired portion of the coding sequence of PRO such as the sequence encoding the extracellular domain of a transmembrane protein or the sequence encoding the mature protein if the protein is extracellular is amplified by PCR with primers complementary to the 5' and 3' regions. The 5' primer may incorporate flanking (selected) restriction enzyme sites. The product is then digested with those selected restriction enzymes and subcloned into the expression vector.

Recombinant baculovirus is generated by co-transfecting the above plasmid and BaculoGoldTM virus DNA (Pharmingen) into *Spodoptera frugiperda* ("Sf9") cells (ATCC CRL 1711) using lipofectin (commercially available from GIBCO-BRL). After 4 - 5 days of incubation at 28°C, the released viruses are harvested and used for further amplifications. Viral infection and protein expression are performed as described by O'Reilley et al., <u>Baculovirus expression vectors</u>: <u>A Laboratory Manual</u>, Oxford: Oxford University Press (1994).

Expressed poly-his tagged PRO can then be purified, for example, by Ni²⁺-chelate affinity chromatography as follows. Extracts are prepared from recombinant virus-infected Sf9 cells as described by Rupert et al., Nature, 362:175-179 (1993). Briefly, Sf9 cells are washed, resuspended in sonication buffer (25 mL Hepes, pH 7.9; 12.5 mM MgCl₂; 0.1 mM EDTA; 10% glycerol; 0.1% NP-40; 0.4 M KCl), and sonicated twice for 20 seconds on ice. The sonicates are cleared by centrifugation, and the supernatant is diluted 50-fold in loading buffer (50 mM phosphate, 300 mM NaCl, 10% glycerol, pH 7.8) and filtered through a 0.45 μm filter. A Ni²⁺-NTA agarose column (commercially available from Qiagen) is prepared

with a bed volume of 5 mL, washed with 25 mL of water and equilibrated with 25 mL of loading buffer. The filtered cell extract is loaded onto the column at 0.5 mL per minute. The column is washed to baseline A₂₈₀ with loading buffer, at which point fraction collection is started. Next, the column is washed with a secondary wash buffer (50 mM phosphate; 300 mM NaCl, 10% glycerol, pH 6.0), which elutes nonspecifically bound protein. After reaching A₂₈₀ baseline again, the column is developed with a 0 to 500 mM Imidazole gradient in the secondary wash buffer. One mL fractions are collected and analyzed by SDS-PAGE and silver staining or Western blot with Ni²⁺-NTA-conjugated to alkaline phosphatase (Qiagen). Fractions containing the eluted His₁₀-tagged PRO are pooled and dialyzed against loading buffer.

Alternatively, purification of the IgG tagged (or Fc tagged) PRO can be performed using known chromatography techniques, including for instance, Protein A or protein G column chromatography.

Many of the PRO polypeptides disclosed herein were successfully expressed as described above.

EXAMPLE 7: Preparation of Antibodies that Bind PRO

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This example illustrates preparation of monoclonal antibodies which can specifically bind PRO.

Techniques for producing the monoclonal antibodies are known in the art and are described, for instance, in Goding, <u>supra</u>. Immunogens that may be employed include purified PRO, fusion proteins containing PRO, and cells expressing recombinant PRO on the cell surface. Selection of the immunogen can be made by the skilled artisan without undue experimentation.

Mice, such as Balb/c, are immunized with the PRO immunogen emulsified in complete Freund's adjuvant and injected subcutaneously or intraperitoneally in an amount from 1-100 micrograms. Alternatively, the immunogen is emulsified in MPL-TDM adjuvant (Ribi Immunochemical Research, Hamilton, MT) and injected into the animal's hind foot pads. The immunized mice are then boosted 10 to 12 days later with additional immunogen emulsified in the selected adjuvant. Thereafter, for several weeks, the mice may also be boosted with additional immunization injections. Serum samples may be periodically obtained from the mice by retro-orbital bleeding for testing in ELISA assays to detect anti-PRO antibodies.

After a suitable antibody titer has been detected, the animals "positive" for antibodies can be injected with a final intravenous injection of PRO. Three to four days later, the mice are sacrificed and the spleen cells are harvested. The spleen cells are then fused (using 35% polyethylene glycol) to a selected murine myeloma cell line such as P3X63AgU.1, available from ATCC, No. CRL 1597. The fusions generate hybridoma cells which can then be plated in 96 well tissue culture plates containing HAT (hypoxanthine, aminopterin, and thymidine) medium to inhibit proliferation of non-fused cells, myeloma hybrids, and spleen cell hybrids.

The hybridoma cells will be screened in an ELISA for reactivity against PRO. Determination of "positive" hybridoma cells secreting the desired monoclonal antibodies against PRO is within the skill in the art.

The positive hybridoma cells can be injected intraperitoneally into syngeneic Balb/c mice to produce ascites containing the anti-PRO monoclonal antibodies. Alternatively, the hybridoma cells can be grown in tissue culture flasks or roller bottles. Purification of the monoclonal antibodies produced in the ascites can be accomplished using ammonium sulfate precipitation, followed by gel exclusion

chromatography. Alternatively, affinity chromatography based upon binding of antibody to protein A or protein G can be employed.

EXAMPLE 8: Purification of PRO Polypeptides Using Specific Antibodies

Native or recombinant PRO polypeptides may be purified by a variety of standard techniques in the art of protein purification. For example, pro-PRO polypeptide, mature PRO polypeptide, or pre-PRO polypeptide is purified by immunoaffinity chromatography using antibodies specific for the PRO polypeptide of interest. In general, an immunoaffinity column is constructed by covalently coupling the anti-PRO polypeptide antibody to an activated chromatographic resin.

Polyclonal immunoglobulins are prepared from immune sera either by precipitation with ammonium sulfate or by purification on immobilized Protein A (Pharmacia LKB Biotechnology, Piscataway, N.J.). Likewise, monoclonal antibodies are prepared from mouse ascites fluid by ammonium sulfate precipitation or chromatography on immobilized Protein A. Partially purified immunoglobulin is covalently attached to a chromatographic resin such as CnBr-activated SEPHAROSETM (Pharmacia LKB Biotechnology). The antibody is coupled to the resin, the resin is blocked, and the derivative resin is washed according to the manufacturer's instructions.

Such an immunoaffinity column is utilized in the purification of PRO polypeptide by preparing a fraction from cells containing PRO polypeptide in a soluble form. This preparation is derived by solubilization of the whole cell or of a subcellular fraction obtained via differential centrifugation by the addition of detergent or by other methods well known in the art. Alternatively, soluble PRO polypeptide containing a signal sequence may be secreted in useful quantity into the medium in which the cells are grown.

A soluble PRO polypeptide-containing preparation is passed over the immunoaffinity column, and the column is washed under conditions that allow the preferential absorbance of PRO polypeptide (e.g., high ionic strength buffers in the presence of detergent). Then, the column is eluted under conditions that disrupt antibody/PRO polypeptide binding (e.g., a low pH buffer such as approximately pH 2-3, or a high concentration of a chaotrope such as urea or thiocyanate ion), and PRO polypeptide is collected.

EXAMPLE 9: Drug Screening

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This invention is particularly useful for screening compounds by using PRO polypeptides or binding fragment thereof in any of a variety of drug screening techniques. The PRO polypeptide or fragment employed in such a test may either be free in solution, affixed to a solid support, borne on a cell surface, or located intracellularly. One method of drug screening utilizes eukaryotic or prokaryotic host cells which are stably transformed with recombinant nucleic acids expressing the PRO polypeptide or fragment. Drugs are screened against such transformed cells in competitive binding assays. Such cells, either in viable or fixed form, can be used for standard binding assays. One may measure, for example, the formation of complexes between PRO polypeptide or a fragment and the agent being tested. Alternatively, one can examine the diminution in complex formation between the PRO polypeptide and its target cell or target receptors caused by the agent being tested.

Thus, the present invention provides methods of screening for drugs or any other agents which can affect a PRO polypeptide-associated disease or disorder. These methods comprise contacting such an agent with an PRO polypeptide or fragment thereof and assaying (I) for the presence of a complex between the agent and the PRO polypeptide or fragment, or (ii) for the presence of a complex between the PRO polypeptide or fragment and the cell, by methods well known in the art. In such competitive binding assays, the PRO polypeptide or fragment is typically labeled. After suitable incubation, free PRO polypeptide or fragment is separated from that present in bound form, and the amount of free or uncomplexed label is a measure of the ability of the particular agent to bind to PRO polypeptide or to interfere with the PRO polypeptide/cell complex.

Another technique for drug screening provides high throughput screening for compounds having suitable binding affinity to a polypeptide and is described in detail in WO 84/03564, published on September 13, 1984. Briefly stated, large numbers of different small peptide test compounds are synthesized on a solid substrate, such as plastic pins or some other surface. As applied to a PRO polypeptide, the peptide test compounds are reacted with PRO polypeptide and washed. Bound PRO polypeptide is detected by methods well known in the art. Purified PRO polypeptide can also be coated directly onto plates for use in the aforementioned drug screening techniques. In addition, non-neutralizing antibodies can be used to capture the peptide and immobilize it on the solid support.

This invention also contemplates the use of competitive drug screening assays in which neutralizing antibodies capable of binding PRO polypeptide specifically compete with a test compound for binding to PRO polypeptide or fragments thereof. In this manner, the antibodies can be used to detect the presence of any peptide which shares one or more antigenic determinants with PRO polypeptide.

EXAMPLE 10: Rational Drug Design

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The goal of rational drug design is to produce structural analogs of biologically active polypeptide of interest (i.e., a PRO polypeptide) or of small molecules with which they interact, e.g., agonists, antagonists, or inhibitors. Any of these examples can be used to fashion drugs which are more active or stable forms of the PRO polypeptide or which enhance or interfere with the function of the PRO polypeptide in vivo (c.f., Hodgson, Bio/Technology, 9: 19-21 (1991)).

In one approach, the three-dimensional structure of the PRO polypeptide, or of a PRO polypeptide-inhibitor complex, is determined by x-ray crystallography, by computer modeling or, most typically, by a combination of the two approaches. Both the shape and charges of the PRO polypeptide must be ascertained to elucidate the structure and to determine active site(s) of the molecule. Less often, useful information regarding the structure of the PRO polypeptide may be gained by modeling based on the structure of homologous proteins. In both cases, relevant structural information is used to design analogous PRO polypeptide-like molecules or to identify efficient inhibitors. Useful examples of rational drug design may include molecules which have improved activity or stability as shown by Braxton and Wells, Biochemistry, 31:7796-7801 (1992) or which act as inhibitors, agonists, or antagonists of native peptides as shown by Athauda et al., J. Biochem., 113:742-746 (1993).

It is also possible to isolate a target-specific antibody, selected by functional assay, as described above, and then to solve its crystal structure. This approach, in principle, yields a pharmacore upon which

subsequent drug design can be based. It is possible to bypass protein crystallography altogether by generating anti-idiotypic antibodies (anti-ids) to a functional, pharmacologically active antibody. As a mirror image of a mirror image, the binding site of the anti-ids would be expected to be an analog of the original receptor. The anti-id could then be used to identify and isolate peptides from banks of chemically or biologically produced peptides. The isolated peptides would then act as the pharmacore.

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By virtue of the present invention, sufficient amounts of the PRO polypeptide may be made available to perform such analytical studies as X-ray crystallography. In addition, knowledge of the PRO polypeptide amino acid sequence provided herein will provide guidance to those employing computer modeling techniques in place of or in addition to x-ray crystallography.

The foregoing written specification is considered to be sufficient to enable one skilled in the art to practice the invention. The present invention is not to be limited in scope by the construct deposited, since the deposited embodiment is intended as a single illustration of certain aspects of the invention and any constructs that are functionally equivalent are within the scope of this invention. The deposit of material herein does not constitute an admission that the written description herein contained is inadequate to enable the practice of any aspect of the invention, including the best mode thereof, nor is it to be construed as limiting the scope of the claims to the specific illustrations that it represents. Indeed, various modifications of the invention in addition to those shown and described herein will become apparent to those skilled in the art from the foregoing description and fall within the scope of the appended claims.